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(FILE 'HOME' ENTERED AT 16:45:22 ON 05 SEP 2006)
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FILE 'HCAPLUS' ENTERED AT 16:45:32 ON 05 SEP 2006

E US2003-734787/APPS

1 SEA ABB=ON PLU=ON US2003-734787/AP T.1 SEL RN

FILE 'REGISTRY' ENTERED AT 16:45:46 ON 05 SEP 2006

L2 6 SEA ABB=ON PLU=ON (158736-49-3/BI OR 252047-40-8/BI OR 263562-55-6/BI OR 338454-52-7/BI OR 57-88-5/BI OR 9028-35-7/BI)

D SCA

FILE 'REGISTRY' ENTERED AT 16:46:12 ON 05 SEP 2006

L3 STR

5 SEA SSS SAM L3 1.4

141 SEA SSS FUL L3 L5

FILE 'HCAPLUS' ENTERED AT 16:47:43 ON 05 SEP 2006

15 SEA ABB=ON PLU=ON L5 L6

FILE 'MEDLINE, EMBASE, BIOSIS' ENTERED AT 16:48:32 ON 05 SEP 2006

0 SEA ABB=ON PLU=ON L5 L7

FILE 'HCAPLUS' ENTERED AT 16:48:40 ON 05 SEP 2006

E ALZHEIM/CT

E E5+ALL

E E2+ALL

23218 SEA ABB=ON PLU=ON "ALZHEIMER'S DISEASE"+PFT,NT/CT  $^{18}$ 

15 SEA ABB=ON PLU=ON L6 OR (L6 AND (L8 OR ALZHEIM?))

L\*\*\* DEL 1 S (L6 AND (L8 OR ALZHEIM?))

L\*\*\* DEL 1 S L10 AND L1

> FILE 'HCAPLUS, MEDLINE, EMBASE, BIOSIS' ENTERED AT 16:50:39 ON 05 SEP 2006 E CANTON T/AU

84 SEA ABB=ON PLU=ON ("CANTON T"/AU OR "CANTON THIERRY"/AU) L10 E PRADIER L/AU

350 SEA ABB=ON PLU=ON ("PRADIER L"/AU OR "PRADIER LAURANT"/AU OR L11 "PRADIER LAURENT"/AU)

E BENAVIDES J/AU

648 SEA ABB=ON PLU=ON ("BENAVIDES J"/AU OR "BENAVIDES J A"/AU OR L12 "BENAVIDES J B"/AU OR "BENAVIDES J E MAURICIO"/AU OR "BENAVIDES J F"/AU OR "BENAVIDES J G"/AU OR "BENAVIDES J I"/AU OR "BENAVIDES J M"/AU OR "BENAVIDES J O"/AU OR "BENAVIDES J R"/AU OR "BENAVIDES JESUS"/AU)

E HEUER H/AU

L13 609 SEA ABB=ON PLU=ON ("HEUER H"/AU OR "HEUER H E"/AU OR "HEUER H G"/AU OR "HEUER H H"/AU OR "HEUER H J"/AU OR "HEUER H O"/AU OR "HEUER H R"/AU OR "HEUER H ROBERT"/AU OR "HEUER H W"/AU OR "HEUER HUBERT"/AU OR "HEUER HUBERT O"/AU OR "HEUER HUBERT OTTO"/AU)

E SCHAEFER H/AU

L\*\*\* DEL 0 S E3, E24, E40E

1853 SEA ABB=ON PLU=ON ("SCHAEFER H"/AU OR "SCHAEFER H L"/AU OR L14"SCHAEFER HANS"/AU OR "SCHAEFER HANS LUDWIG"/AU OR "SCHAEFER HANS LUDWING"/AU)

55 SEA ABB=ON PLU=ON (L10 AND (L11 OR L12 OR L13 OR L14)) OR L15

		(L11 AND (L12 OR L13 OR L14)) OR (L12 AND (L13 OR L14)) OR
		(L13 AND L14)
L16	3484	SEA ABB=ON PLU=ON (L10 OR L11 OR L12 OR L13 OR L14)
L17	4	SEA ABB=ON PLU=ON L16 AND L5
L18	219	SEA ABB=ON PLU=ON L16 AND ALZHEIM?
L19	· 13	SEA ABB=ON PLU=ON L16 AND REUP?
L20	0	SEA ABB=ON PLU=ON L18 AND L19
L21	54	SEA ABB=ON PLU=ON L16 AND INTEST?
L22	6	SEA ABB=ON PLU=ON L16 AND INTEST? (5A) ?INHIB?
L23	519	SEA ABB=ON PLU=ON L16 AND INHIB?
L24	69	SEA ABB=ON PLU=ON L23 AND (INTEST? OR HMG? OR REDUCTAS? OR
		CHOLEST? OR APP OR SECRETAS? OR REUP? OR BILIAR?)
L25	121	SEA ABB=ON PLU=ON L24 OR L22 OR L17 OR L15
L26	65	DUP REM L25 (56 DUPLICATES REMOVED)
		ANSWERS '1-50' FROM FILE HCAPLUS
		ANSWERS '51-52' FROM FILE MEDLINE
		ANSWER '53' FROM FILE EMBASE
		ANSWERS '54-65' FROM FILE BIOSIS

=> fil hcap

FILE 'HCAPLUS' ENTERED AT 16:59:01 ON 05 SEP 2006
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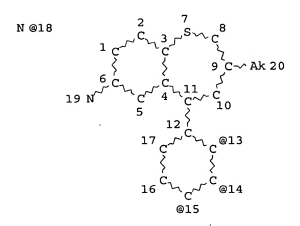
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FILE COVERS 1907 - 5 Sep 2006 VOL 145 ISS 11 FILE LAST UPDATED: 4 Sep 2006 (20060904/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

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VPA 18-13/14/15 U NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

**GRAPH ATTRIBUTES:** 

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 20

STEREO ATTRIBUTES: NONE

L5 141 SEA FILE=REGISTRY SSS FUL L3

L6 15 SEA FILE=HCAPLUS ABB=ON PLU=ON

L8 23218 SEA FILE=HCAPLUS ABB=ON PLU=ON "ALZHEIMER'S DISEASE"+PFT,NT/C

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L9 15 SEA FILE=HCAPLUS ABB=ON PLU=ON L6 OR (L6 AND (L8 OR ALZHEIM?) )

=> d 19 ibib abs hitind hitstr 1-15

ANSWER 1 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2005:979630 HCAPLUS

DOCUMENT NUMBER:

143:286456

TITLE:

Preparation of benzothiazepine and benzothiepine

compounds for prevention and treatment of hyperlipemia

INVENTOR(S):

Sasahara, Takehiko; Mohri, Mitsunobu; Kasahara,

Kenichi

PATENT ASSIGNEE(S):

Asahi Kasei Pharma Corporation, Japan

PCT Int. Appl., 551 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

Japanese

1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.						KIN	<b>D</b> :	DATE			APPL	ICAT:	DATE					
							-											
WO 2005082874					<b>A1</b>		2005	0909	1	WO 2	005-i		20050225					
		W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	ΚP,	KR,	ΚZ,	LC,
			LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,

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NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

PRIORITY APPLN. INFO::

JP 2004-128992 A 20040227

OTHER SOURCE(S):

MARPAT 143:286456
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\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

Benzothiazepine or benzothiepine compds. represented by the following AB general formula (I) which have a thioamide bond and a quaternary ammonium substituent [R1a, R2a = C1-10 alkyl, C2-10 alkenyl, C2-10 alkynyl; ma = an integer of 0-4; Rx = halo, NO2, NH2, cyano, HO, CO2H, CONH2, SO3H, C1-5 alkylated NH2, each (un)substituted C1-10 alkyl, C2-10 alkenyl, or C2-10 alkynyl; a combination of (A1, A2, A3) = (CH2, NH, CH), (CH2, CH(OH), CH), (NH, CH(OH), CH), or (CH2, CH2, N); Y = NHC(S), NHC(S)NH, NHC(S)O; Za-(N+R5aR6aR7a)n = C2-10 alkyl or alkenyl substituted by n number of (N+R5aR6aR7a) with ≥1 CH2 of Za optionally being replaced with (un) substituted phenylene or O; n = 1,2; N+R5aR6aR7a is selected from (1) R5a, R6a, R7a = each (un) substituted C1-10 alkyl, C2-10 alkenyl, or C2-10 alkynyl with ≥1 CH2 optionally being replaced with phenylene, thienylene, furylene, cyclohexylene, cyclopentylene, O, S, CO2, NHCO, (un) substituted NH, or ammonium ion, (2) C4-9 mono- or bicyclic ring containing ammonium N atom with at least one of the ring C atoms being replaced with O, N, or S, and (3) (un)substituted pyridinium, quinolinium, or isoquinolinium; X- = a counter ion] are prepared These compds. are useful as remedies and preventives for hyperlipemia, arteriosclerosis, syndrome X, and other coronary artery diseases and as cholesterol-lowering agents. They are also useful as remedies and preventives for liver disorders accompanying bile stasis, in particular, primary biliary cirrhosis, and primary sclerotic cholangitis, and obesity, fatty liver, and fatty hepatitis. Thus, 95 mg 1-(4-isocyanatobenzyl)-1azoniabicyclo[2.2.2]octane bromide (preparation given) was added to a solution of

116 mg 5-(3-aminophenyl)-3,3-dibutyl-7-dimethylamino-4-hydroxy-2,3,4,5-tetrahydro-1-benzothiepin-1,1-dioxide in 3 mL CHCl3 and stirred at 50° for 2 h to give 190 mg -2,3,4,5-tetrahydro-1-benzothiepin-1,1-dioxide derivative (II). II at 0.3 mg/kg twice a day for 5 days lowered a total blood level of LDL and VLDL by 49% in rats on 5 day diet of a feed containing 0.5% cholesterol and 0.5% bile acid.

IC ICM C07D281-10

ICS A61K031-22; A61K031-366; A61K031-38; A61K031-381; A61K031-395; A61K031-4025; A61K031-428; A61K031-439; A61K031-4995; A61K031-5375; A61K031-541; A61K031-554; A61P001-16; A61P003-04; A61P003-06; A61P009-10; C07D337-08; C07D409-12; C07D453-02

CC 28-22 (Heterocyclic Compounds (More Than One Hetero Atom)) Section cross-reference(s): 1, 27

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670277-37-9P

670277-35-7P

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300350-08-7P

393856-01-4P

300350-09-8P

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RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
   (preparation of benzothiazepine and benzothiepine compds. for prevention and
   treatment of hyperlipemia and lowering cholesterol)
727-93-5P, 4-Fluoro-2-(4-methoxybenzoyl)phenol
                                                  729-39-5P,
4-Methoxybenzoic acid 4-fluorophenyl ester
                                              6141-45-3P, Methyl
2-amino-2-butylhexanoate
                            31520-33-9P, 5-Bromopentyl isothiocyanate
31520-34-0P, 6-Bromohexyl isothiocyanate
                                           40216-70-4P, Methyl
2-(benzylideneamino)hexanoate
                                 73674-09-6P, 8-Bromooctanoyl chloride
90331-59-2P, 4-(2-Bromoethyl)phenyl isothiocyanate
                                                       105732-43-2P,
2-Aminoheptanoic acid methyl ester hydrochloride
                                                    155863-31-3P
155863-32-4P, 4-(Bromomethyl)phenyl isothiocyanate
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197378-15-7P
                               197378-18-0P
                                                              288161-81-9P,
2-Amino-2-butylhexanol
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300350-10-1P

393855-85-1P

670278-59-8P, Methyl 2-(benzylideneamino)-2-butylhexanoate

393855-98-6P

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670278-60-1P, 2-[(2-Amino-2-butylhexyl)thio]-5-fluorobenzophenone
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3,3-Dibutyl-2,3-dihydro-7-fluoro-5-(4-methoxyphenyl)-1,4-benzothiazepine-
1,1-dioxide
              670278-72-5P, 3,3-Dibutyl-2,3-dihydro-7-dimethylamino-5-(4-
methoxyphenyl)-1,4-benzothiazepine-1,1-dioxide
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RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
   (preparation of benzothiazepine and benzothiepine compds. for prevention and
   treatment of hyperlipemia and lowering cholesterol)
864348-68-5P 864348-69-6P 864348-70-9P
864348-71-0P 864348-72-1P 864348-73-2P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
   (preparation of benzothiazepine and benzothiepine compds. for prevention and
   treatment of hyperlipemia and lowering cholesterol)
864348-68-5 HCAPLUS
4-Aza-1-azoniabicyclo[2.2.2]octane, 1-[[4-[[[[3-[3,3-dibuty]-7-
(dimethylamino) -2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-
```

yl]phenyl]amino]thioxomethyl]amino]phenyl]methyl]-, bromide (9CI) (CA

IT

RN

CN

INDEX NAME)

RN 864348-69-6 HCAPLUS
CN 4-Aza-1-azoniabicyclo[2.2.2]octane, 1-[[4-[[[[3-[3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]thioxomethyl]amino]-3-fluorophenyl]methyl]-, bromide (9CI) (CA INDEX NAME)

RN 864348-70-9 HCAPLUS

CN 1-Azoniabicyclo[2.2.2]octane, 1-[[4-[[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]thioxomethyl]amino]phenyl]methyl]-4-phenyl-, bromide, rel-(9CI) (CA INDEX NAME)

PAGE 2-A

• Br-

RN 864348-71-0 HCAPLUS
CN Pyridinium, 1-[[4-[[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]thioxomethyl]amino]phenyl]methyl]-4-(1,1-dimethylethyl)-,bromide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

● Br -

RN 864348-72-1 HCAPLUS

CN Pyridinium, 1-[[4-[[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]thioxomethyl]amino]phenyl]methyl]-3-(3-methoxy-3-oxopropyl)-, bromide, rel- (9CI) (CA INDEX NAME)

## PAGE 1-A

PAGE 2-A

Br -

RN 864348-73-2 HCAPLUS

CN

Pyridinium, 1-[[4-[[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]thioxomethyl]amino]phenyl]methyl]-2-propyl-, bromide, rel-(9CI) (CA INDEX NAME)

Relative stereochemistry.

## PAGE 1-A

PAGE 2-A

● Br~

IT 864350-14-1P 864350-15-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of benzothiazepine and benzothiepine compds. for prevention and treatment of hyperlipemia and lowering cholesterol)

RN 864350-14-1 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-5-(3-nitrophenyl)-, 1,1-dioxide (9CI) (CA INDEX NAME)

RN 864350-15-2 HCAPLUS

CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-, 1,1-dioxide (9CI) (CA INDEX NAME)

REFERENCE COUNT:

14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 2 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2005:704248 HCAPLUS

DOCUMENT NUMBER:

143:347042

TITLE:

Discovery of Potent, Nonsystemic Apical

Sodium-Codependent Bile Acid Transporter Inhibitors

(Part 2)

AUTHOR(S):

Huang, Horng-Chih; Tremont, Samuel J.; Lee, Len F.; Keller, Bradley T.; Carpenter, Andrew J.; Wang, Ching-Cheng; Banerjee, Shyamal C.; Both, Scott R.;

Fletcher, Theresa; Garland, Danny J.; Huang, Wei; Jones, Claude; Koeller, Kevin J.; Kolodziej, Steve A.; Li, James; Manning, Robert E.; Mahoney, Matthew W.; Miller, Raymond E.; Mischke, Deborah A.; Rath, Nigam P.; Reinhard, Emily J.; Tollefson, Michael B.; Vernier, William F.; Wagner, Grace M.; Rapp, Steve R.; Beaudry, Judy; Glenn, Kevin; Regina, Karen; Schuh, Joe R.; Smith, Mark E.; Trivedi, Jay S.; Reitz, David B. Discovery Chemistry and Department of Cardiovascular Disease, Pharmacia, Chesterfield, MO, 63017, USA Journal of Medicinal Chemistry (2005), 48(18),

5853-5868

CODEN: JMCMAR; ISSN: 0022-2623 American Chemical Society

Journal English

LANGUAGE: English
OTHER SOURCE(S): CASREACT 143:347042

Ι

GI

SOURCE:

PUBLISHER:

DOCUMENT TYPE:

CORPORATE SOURCE:

AB Since the primary site for active bile acid reabsorption is via apical sodium-codependent bile acid transporter (ASBT), which is localized on the luminal surface of the distal ileum, a nonsystemic inhibitor would be desirable to minimize or eliminate potential systemic side effects of an absorbed drug. To ensure bioequivalency and product stability, it was also essential that a nonhygroscopic inhibitor in its most stable crystalline form was identified. A series of benzothiepins I [R = Ph, 4-HOC6H4, 4-(Me2NCH2CH2)C6H4, 1-naphthyl, 2-thienyl, 3-pyridyl, etc.] was prepared to refine the structure-activity relationship of the substituted Ph ring at the 5-position of benzothiepin ring and to identify potent, crystalline, nonhygroscopic, and efficacious ASBT inhibitors with low systemic exposure.

CC 27-21 (Heterocyclic Compounds (One Hetero Atom))
Section cross-reference(s): 1

IT 228113-65-3P 289038-80-8P 865593-98-2P 865593-99-3P 865594-02-1P 865594-03-2P **865594-07-6P 865594-11-2P** 865594-17-8P 865594-23-6P 865594-31-6P 865594-36-1P

RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent)

(preparation of (aryl) (hydroxy) tetrahydrobenzothiepins as nonsystemic apical sodium-codependent bile acid transporter inhibitors)

IT 197373-42-5P 197373-43-6P 197373-48-1P 197373-51-6P 197373-56-1P 197373-57-2P 197373-64-1P 197374-40-6P 197374-45-1P 197374-48-4P 197374-67-7P 197374-72-4P 197374-92-8P 197374-96-2P 197375-70-5P 197376-07-1P 197376-11-7P 197376-19-5P 197376-22-0P 197376-34-4P 197376-46-8P 197376-58-2P 197376-67-3P 197376-76-4P 197377-03-0P 197377-38-1P 197377-47-2P 197377-51-8P 197377-62-1P

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289037-76-9P
               865593-97-1P
                               865594-00-9P
                                              865594-05-4P
                                                              865594-06-5P
865594-09-8P 865594-12-3P 865594-14-5P
865594-15-6P
               865594-16-7P
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                               865594-25-8P
                                              865594-26-9P
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865594-22-5P
865594-28-1P
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865594-34-9P
865594-40-7P
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                                              865594-50-9P
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                                                              865594-59-8P
865594-53-2P
865594-60-1P
               865594-61-2P
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RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation of (aryl) (hydroxy)tetrahydrobenzothiepins as nonsystemic apical sodium-codependent bile acid transporter inhibitors)

IT 197378-13-5P 197378-15-7P 197378-16-8P 197378-18-0P 289038-53-5P 289038-61-5P 865593-96-0P 865594-01-0P 865594-04-3P 865594-08-7P 865594-10-1P 865594-56-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of (aryl) (hydroxy) tetrahydrobenzothiepins as nonsystemic apical sodium-codependent bile acid transporter inhibitors)

IT 865594-07-6P 865594-11-2P

RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent)

(preparation of (aryl) (hydroxy) tetrahydrobenzothiepins as nonsystemic apical sodium-codependent bile acid transporter inhibitors)

RN 865594-07-6 HCAPLUS

CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RN 865594-11-2 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-5-[3-(4-pyridinylamino)phenyl]-, 1,1-dioxide, (4R,5R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

197373-51-6P 865594-09-8P 865594-12-3P IT

865594-14-5P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL

(Biological study); PREP (Preparation) (preparation of (aryl) (hydroxy) tetrahydrobenzothiepins as nonsystemic apical sodium-codependent bile acid transporter inhibitors)

197373-51-6 HCAPLUS RN

1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3,3-dibutyl-7-(dimethylamino)-CN2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 865594-09-8 HCAPLUS

Glycine, N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-CN hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]glycyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RN 865594-12-3 HCAPLUS

CN Pyridinium, 4-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-1-methyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

• I-

RN 865594-14-5 HCAPLUS

CN 1-Pentanaminium, 5-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-5-oxo-, salt with trifluoroacetic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 865594-13-4 CMF C37 H60 N3 O4 S Absolute stereochemistry. Rotation (+).

CM 2

CRN 14477-72-6 CMF C2 F3 O2

IT 865594-08-7P 865594-10-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of (aryl) (hydroxy) tetrahydrobenzothiepins as nonsystemic apical sodium-codependent bile acid transporter inhibitors)

RN 865594-08-7 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-5-[3-[(diphenylmethylene)amino]phenyl]-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 865594-10-1 HCAPLUS

CN Glycine, N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]glycyl-, ethyl ester (9CI) (CA INDEX NAME)

## Absolute stereochemistry.

REFERENCE COUNT: 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 3 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:492306 HCAPLUS

DOCUMENT NUMBER: 141:17641

TITLE: Methods and compositions for the prevention and

treatment of Alzheimer's disease with intestinal bile acid reuptake inhibitors

PATENT ASSIGNEE(S): Aventis Pharma SA, Fr. SOURCE: Fr. Demande, 25 pp.

CODEN: FRXXBL

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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KIND
                               DATE
                                         APPLICATION NO.
                                                                 DATE
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    FR 2848452
                        A1
                               20040618 FR 2002-15722
                                                                 20021212
    CA 2507945
                        AA
                               20040729 CA 2003-2507945
    WO 2004062652
                        A1
                               20040729
                                        WO 2003-FR3654
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE,
            GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
            LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ,
            OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM,
            TN, TR, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW
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            FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
    AU 2003296802
                         Α1
                               20040810
                                        AU 2003-296802
                                                                 20031210
                               20050914
                                          EP 2003-815109
    EP 1572174
                         A1
                                                                 20031210
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
    BR 2003017280
                        Α
                               20051108
                                          BR 2003-17280
                                                                 20031210
    CN 1726016
                        Α
                               20060125 CN 2003-80105972
                                                                 20031210
                                         JP 2004-566119
    JP 2006514063
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                               20060427
                                                                 20031210
                               20040715 US 2003-734787
    US 2004138145
                        A1
                                                                 20031212
    NO 2005003341
                        Α
                               20050907
                                          NO 2005-3341
                                                                 20050708
PRIORITY APPLN. INFO.:
                                          FR 2002-15722
                                                             A 20021212
                                          US 2003-455354P P 20030317
WO 2003-FR3654 W 20031210
OTHER SOURCE(S):
                        MARPAT 141:17641
    The invention describe the application of the intestinal biliary acid
    reuptake inhibitors for the prevention and the treatment of
    Alzheimer's disease, alone or in conjunction with an HMG-COA
    reductase inhibitor , a cholesterol uptake inhibitor, a cholesterol
    synthesis inhibitor or an inhibitor of APP secretases.
IC
    ICM A61K031-444
    ICS A61K031-38; A61P025-28
```

- CC 1-11 (Pharmacology)
- ST bile acid reuptake inhibitors intestine Alzheimers disease treatment prevention
- IT Intestine

(biliary acid reuptake; methods and compns. for prevention and treatment of Alzheimer's disease with intestinal bile acid reuptake inhibitors)

IT Alzheimer's disease

Anti-Alzheimer's agents

Human

(methods and compns. for prevention and treatment of Alzheimer 's disease with intestinal bile acid reuptake inhibitors)

IT Bile acids

> RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(reuptake inhibitors; methods and compns. for prevention and treatment of Alzheimer's disease with intestinal bile acid reuptake inhibitors)

IT Biological transport

> (reuptake, bile acid, inhibitors of; methods and compns. for prevention and treatment of Alzheimer's disease with intestinal bile acid reuptake inhibitors)

IT Biological transport

(uptake, cholesterol, inhibitors of, in conjunction with treatment; methods and compns. for prevention and treatment of Alzheimer 's disease with intestinal bile acid reuptake inhibitors) TT 9028-35-7, HMG-CoA reductase 158736-49-3, β-Secretase 338454-52-7, γ Secretase RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (inhibitor, in conjunction with treatment; methods and compns. for prevention and treatment of Alzheimer's disease with intestinal bile acid reuptake inhibitors) 263562-55-6 IT 252047-40-8 RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (methods and compns. for prevention and treatment of Alzheimer 's disease with intestinal bile acid reuptake inhibitors) 57-88-5, Cholesterol, biological studies IT RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(Biological study); USES (Uses)
(uptake and synthesis inhibitors, in conjunction with treatment;

(uptake and synthesis inhibitors, in conjunction with treatment; methods and compns. for prevention and treatment of Alzheimer 's disease with intestinal bile acid reuptake inhibitors) 252047-40-8

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(methods and compns. for prevention and treatment of Alzheimer
's disease with intestinal bile acid reuptake inhibitors)

RN 252047-40-8 HCAPLUS

IT

CN D-Glucitol, 1-[[5-[[3-[(3S,4R,5R)-3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-5-oxopentyl]amino]-1-deoxy-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 4 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:60147 HCAPLUS

DOCUMENT NUMBER: 140:111291

TITLE: Preparation of substituted 5-aryl-benzothiepines as

FI

20010409

20020215

ileal bile acid transport and taurocholate uptake

inhibitors

Lee, Len F.; Banerjee, Shyamal C.; Huang, Horng Chih; INVENTOR(S):

Li, Jinglin J.; Miller, Raymond E.; Reitz, David B.;

Tremont, Sanuel J.

PATENT ASSIGNEE(S):

G.D. Searle and Co., USA

SOURCE:

U.S. Pat. Appl. Publ., 235 pp., Cont.-in-part of U.S.

US 2001-828968

US 2002-76091

Ser. No. 831,284.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

9

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.						)	DATE	3	7	LICA	DATE								
			<b>-</b> -			-											·		
US	2004	0148	03		A1		2004	10122	τ	JS 2	2002	-682	97			20	020	208	
US	6784	201			B2		2004	10831											
CA	2506	703			AA		1997	70918	(	CA :	1997-	-2506	5703	3		19	970	311	
ΕP	1440	972			A1		2004	10728	1	EP 2	2004	-1008	8 8			19	970	311	
	R:	AT,	BE,	CH,	DE,	DK,	, ES	FR,	GB,	GR,	, IT	, LI	, LU	J,	NL,	SE,	PT,	ΙE,	
ΑU	7612	49			B2		2003	30529	I	AU 2	2000-	-533	94			20	000	816	

20020131

US 2002013476 A1 US 6387924 B2 20020514 US 2003171426 **A**1 20030911 US 6642268 B2 20031104 A1 20041014

US 2004204478 PRIORITY APPLN. INFO.: US 2004-830125 20040423 US 1994-305526 B2 19940913 B1 19950821 US 1995-517051 P 19960311

US 1996-13119P A2 19970311 US 1997-816065 A2 19970331 US 1997-831284 A3 20010409 US 2001-828968 A3 19970311 AU 1997-23266 CA 1997-2248586 A3 19970311 EP 1997-915976 A3 19970311 Ρ US 1997-40660P 19970311

Р 19971219 US 1997-68170P A2 19980702 US 1998-109551 A1 19990324 US 1999-275463 US 1999-443403 A1 19991119

A3 20000929 US 2000-676466 A3 20020208 US 2002-68297

OTHER SOURCE(S):

MARPAT 140:111291

GI

$$(R?) q \qquad \qquad (R?) q \qquad \qquad (R.) q \qquad$$

MeO 
$$\stackrel{\circ}{\underset{\text{NeO}}{\bigvee}}$$
 Et  $\stackrel{\circ}{\underset{\text{MeO}}{\bigvee}}$  Bu  $\stackrel{\circ}{\underset{\text{Ph}}{\bigvee}}$  OH III

AB The title compds. (I) [wherein q = 1-4; n = 0-2; R1, R2 = H, (un) substituted (halo) alkyl, alkenyl, alkynyl, alkylaryl, arylalkyl, alkoxy(alkyl), dialkylamino, alkylthio, (polyalkyl)aryl, or cycloalkyl; or R1 and R2 taken together with the atoms to which they are attached = cycloalkyl; R3, R4 = H, alkyl, alkenyl, alkynyl, acyloxy, aryl, heterocyclyl, OR9, NR9R10, SR9, S(O)R9, SO2R9, or SO3R9; R9, R10 = H, (cyclo)alkyl, alkenyl, alkynyl, aryl(alkyl), acyl, heterocyclyl, or ammoniumalkyl; or R3 and R4 together = :0, :NOR11, :S, :NNR11R12, :NR9, or :CR11R12; R11, R12 = H, (cyclo)alkyl, alkenyl, alkynyl, aryl(alkyl), heterocyclyl, carboxylalkyl, carboalkoxyalkyl, cyanoalkyl, OR9, NR9R10, SR9, S(0)R9, SO2R9, SO3R9, CO2R9, CN, halo, oxo, or CONR9R10; R5, R6 = H, alkyl, aryl, etc.; R7, R8 = H, alkyl; Rx = H, (un)substituted (cyclo)alkyl, alkenyl, alkynyl, polyalkyl, acyloxy, aryl(alkyl), halo(alkyl), (quaternary) heterocyclyl, (quaternary) heteroaryl, polyether, alkoxy, amino, alkylthio, NO2, carboxy, carbamido, etc.] were prepared for the prophylaxis and treatment of hyperlipidemic conditions, such as those associated with atherosclerosis or hypercholesterolemia. Thus, KOBu-t was added to a solution of 2-((2-benzyl-5methoxyphenylsulfonyl)methyl)-2-ethylhexanal (preparation given) and dry THF cooled to -1.6°C to give, after workup, II and III (96% combined yield). The isomers were separated upon recrystn. II inhibited IBAT-mediated uptake of [14C]-taurocholate in H14 cells with an IC50 of 0.1  $\mu M$  and reduced serum cholesterol from 143 mg (7%) to 126 mg (2%) compared to control in cholesterol-fed hamsters in a 14-day test. In vitro taurocholate uptake assay data are included for nearly 600 compds. of the invention.

IC ICM C07D337-16 ICS A61K031-38

INCL 514431000; 549012000

CC 27-21 (Heterocyclic Compounds (One Hetero Atom))
 Section cross-reference(s): 1, 63

IT 178678-22-3P 178678-23-4P 178678-24-5P 178678-25-6P 178678-26-7P 178678-33-6P 178678-27-8P 178678-29-0P 178678-34-7P 178678-37-0P 178678-50-7P 178678-51-8P 178678-57-4P 178678-46-1P 178678-49-4P 178678-58-5P 178678-59-6P 178897-97-7P 178897-98-8P 178898-00-5P 197372-71-7P 197372-76-2P 197372-77-3P 178898-05-0P 197372-67-1P 197373-42-5P 197373-43-6P 197373-44-7P 197373-47-0P 197372-78-4P 197373-49-2P 197373-50-5P 197373-51-6P 197373-55-0P

197375-48-7P

197373-58-3P

197373-56-1P

197373-57-2P

197375-49-8P

RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as ileal bile acid transport and taurocholate uptake inhibitors) IT 178678-28-9P 178678-30-3P 178678-31-4P 178678-35-8P 178678-36-9P 178678-39-2P 178678-40-5P 178678-41-6P 178678-42-7P 178678-43-8P 178678-44-9P 178678-45-0P 178678-47-2P 178678-48-3P 178678-52-9P 178678-54-1P 178678-53-0P 178897-95-5P 178897-96-6P 178897-99-9P 178898-01-6P 178898-02-7P 178898-03-8P 178898-04-9P 197372-66-0P 197372-70-6P 197372-69-3P 197372-72-8P 197372-73-9P 197372-74-0P 197372-75-1P 197372-79-5P 197372-80-8P 197372-81-9P 197372-82-0P 197372-83-1P 197372-84-2P 197372-85-3P 197372-86-4P 197372-87-5P 197372-88-6P 197372-89-7P 197372-90-0P 197372-91-1P 197372-92-2P 197372-93-3P 197372-94-4P 197372-95-5P 197372-96-6P 197372-97-7P 197372-98-8P 197372-99-9P 197373-00-5P 197373-01-6P 197373-02-7P 197373-03-8P 197373-04-9P 197373-05-0P 197373-06-1P 197373-07-2P 197373-08-3P 197373-09-4P 197373-10-7P 197373-11-8P 197373-12-9P 197373-16-3P 197373-18-5P 197373-13-0P 197373-14-1P 197373-17-4P 197373-19-6P 197373-20-9P 197373-24-3P 197373-22-1P 197373-25-4P 197373-26-5P 197373-27-6P 197373-28-7P 197373-29-8P 197373-30-1P 197373-36-7P **197373-37-8**P 197373-35-6P 197373-38-9P 197373-39-0P 197373-40-3P 197373-41-4P 197373-45-8P 197373-48-1P 197373-54-9P 197373-59-4P 197373-60-7P 197373-61-8P 197373-62-9P 197373-63-0P 197373-64-1P 197373-66-3P 197373-67-4P 197373-68-5P 197373-69-6P 197373~70-9P 197373-71-0P 197373-72-1P 197373-73-2P 197373-75-4P 197373-78-7P 197373-76-5P 197373-77-6P 197373-80-1P 197373-79-8P 197373-81-2P 197373-83-4P 197373-85-6P 197373-87-8P 197373-90-3P · 197373-93-6P 197373-95-8P 197373-97-0P 197373-99-2P 197374-00-8P 197374-01-9P 197374-02-0P 197374-03-1P 197374-04-2P 197374-09-7P 197374-06-4P 197374-08-6P 197374-10-0P 197374-11-1P 197374-13-3P 197374-14-4P 197374-16-6P 197374~17-7P 197374-18-8P 197374-19-9P 197374-20-2P 197374-21-3P 197374-22-4P 197374-24-6P 197374-25-7P 197374-26-8P 197374-27-9P 197374-29-1P 197374-30-4P 197374-31-5P 197374-32-6P 197374-34-8P 197374-35-9P 197374-37-1P 197374-38-2P 197374-39-3P 197374-40-6P 197374-41-7P 197374-43-9P 197374-44-0P 197374-45-1P 197374-46-2P 197374-47-3P 197374-49-5P 197374-48-4P 197374-50-8P 197374-51-9P 197374-52-0P 197374-53-1P 197374-54-2P 197374-55-3P 197374-56-4P 197374-58-6P **197374-59-7P** 197374-57-5P 197374-60-0P 197374-62-2P 197374-63-3P 197374-64-4P 197374-65-5P 197374-66-6P 197374-67-7P 197374-68-8P 197374-69-9P 197374-71-3P 197374-72-4P 197374-76-8P 197374-73-5P 197374-74-6P 197374-75-7P 197374-77-9P 197374-78-0P 197374-79-1P 197374-80-4P 197374-81-5P 197374-82-6P 197374-83-7P 197374-84-8P 197374-85-9P 197374-86-0P 197374-87-1P 197374-88-2P 197374-89-3P 197374-90-6P 197374-91-7P 197374-92-8P 197374-93-9P 197374-94-0P 197374-95-1P 197374-96-2P 197374-97-3P 197374-98-4P 197374-99-5P 197375-00-1P 197375-02-3P 197375-01-2P 197375-03-4P 197375-04-5P 197375-05-6P 197375-06-7P 197375-07-8P 197375-08-9P 197375-09-0P 197375-10-3P 197375-11-4P 197375-12-5P 197375-13-6P 197375-14-7P 197375-15-8P 197375-16-9P 197375-17-0P 197375-20-5P 197375-22-7P 197375-23-8P 197375-24-9P 197375-25-0P 197375-26-1P 197375-28-3P 197375-30-7P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by

```
cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as
        ileal bile acid transport and taurocholate uptake inhibitors)
IT
                    197375-34-1P
                                    197375-39-6P
                                                   197375-42-1P
                                                                   197375-44-3P
     197375-32-9P
     197375-52-3P
                    197375-57-8P
                                    197375-60-3P
                                                   197375-63-6P.
                                                                   197375-66-9P
     197375-68-1P
                    197375-70-5P
                                    197375-72-7P
                                                   197375-74-9P
                                                                   197375-75-0P
     197375-80-7P
                    197375-82-9P
                                    197375-84-1P
                                                   197375-86-3P
                                                                   197375-89-6P
                    197375-94-3P 197375-96-5P
                                                 197375-98-7P
     197375-93-2P
                                                                   197376-07-1P
     197376-00-4P
                    197376-02-6P
                                    197376-04-8P
                                                   197376-06-0P
                    197376-09-3P
                                    197376-10-6P
                                                   197376-11-7P
                                                                   197376-12-8P
     197376-08-2P
                    197376-14-0P
                                    197376-15-1P
                                                   197376-17-3P
                                                                   197376-18-4P
     197376-13-9P
                    197376-21-9P
                                    197376-22-0P
                                                   197376-25-3P
                                                                   197376-31-1P
     197376-19-5P
                    197376-34-4P
                                    197376-36-6P
                                                   197376-38-8P
                                                                   197376-40-2P
     197376-32-2P
     197376-42-4P
                    197376-46-8P
                                    197376-49-1P
                                                   197376-52-6P
                    197376-58-2P
                                    197376-61-7P
                                                   197376-64-0P
     197376-55-9P
                    197376-69-5P
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                                                   197376-74-2P
                                                                   197376-75-3P
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                    197376-77-5P
                                    197376-78-6P
                                                   197376-79-7P
                                                                   197376-81-1P
     197376-76-4P
                                                                   197376-86-6P
     197376-82-2P
                    197376-83-3P
                                    197376-84-4P
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                    197376-89-9P
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     197376-95-7P
                    197376-97-9P
                                    197376-99-1P
                                                   197377-00-7P
                                                                   197377-02-9P
                    197377-05-2P
                                    197377-09-6P
                                                   197377-10-9P
                                                                   197377-11-0P
     197377-03-0P
                                    197377-16-5P
                                                   197377-17-6P
                                                                   197377-18-7P
     197377-12-1P
                    197377-14-3P
                    197377-20-1P
                                    197377-21-2P
                                                   197377-22-3P
                                                                   197377-23-4P
     197377-19-8P
                    197377-25-6P
                                    197377-26-7P
                                                   197377-27-8P
                                                                   197377-28-9P
     197377-24-5P
     197377-29-0P
                    197377-30-3P
                                    197377-31-4P
                                                   197377-32-5P
                                                                   197377-33-6P
                    197377-35-8P
                                    197377-36-9P
                                                   197377-37-0P
                                                                   197377-38-1P
     197377-34-7P
                    197377-40-5P
                                    197377-42-7P
                                                   197377-43-8P
                                                                   197377-45-0P
     197377-39-2P
                    197377-47-2P
                                    197377-48-3P
                                                   197377-49-4P
                                                                   197377-50-7P
     197377-46-1P
                    197377-53-0P
                                    197377-54-1P
                                                   197377-55-2P
                                                                   197377-57-4P
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     197377-58-5P
                    197377-60-9P
                                    197377-61-0P
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                    197377-65-4P
                                    197377-66-5P
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     197377-70-1P
                    197377-71-2P
                                    197377-72-3P
                                                   197377-73-4P
                                                                   197377-74-5P
                    197377-76-7P
                                    197377-77-8P
                                                   197377-78-9P
                                                                   197377-79-0P
     197377-75-6P
                    197377-82-5P
                                    197377-83-6P
                                                   197377-84-7P
                                                                   197377-85-8P
     197377-81-4P
                    197377-90-5P
                                    197377-94-9P
                                                   197377-96-1P
                                                                   197377-98-3P
     197377-86-9P
     197384-36-4P
                    197384-39-7P
                                    197390-49-1P
                                                   197390-68-4P
                                    213312-99-3P
                                                   213313-15-6P
                                                                   213313-34-9P
     213312-50-6P
                    213312-80-2P
                                    289037-53-2P
                                                   289037-54-3P
                                                                   289037-55-4P
     213386-72-2P
                    228113-66-4P
                    289037-57-6P
                                    289037-58-7P
                                                   289037-59-8P
                                                                   289037-60-1P
     289037-56-5P
                    289037-62-3P
                                    289037-64-5P
                                                   289037-65-6P
                                                                   289037-67-8P
     289037-61-2P
     289037-68-9P
                    289037-70-3P
                                    289037-72-5P
                                                   289037-74-7P
                                                                   289037-75-8P
                    289037-77-0P
                                    289037-78-1P
                                                   289037-79-2P
                                                                   289037-80-5P
     289037-76-9P
     289037-81-6P
                    289037-82-7P
                                    289037-83-8P
                                                   289037-84-9P
                                                                   289037-85-0P
     289037-86-1P
                    289037-87-2P
                                    289037-88-3P
                                                   289037-90-7P
                                                                   289037-91-8P
                    289037-93-0P
                                    289037-94-1P
                                                   289037-95-2P
                                                                   289038-00-2P
     289037-92-9P
                                    289038-03-5P
                                                   289038-04-6P
                                                                   289038-05-7P
     289038-01-3P
                    289038-02-4P
     289038-06-8P
                    289038-07-9P
                                    289038-09-1P
                                                   289038-11-5P
                                                                   289038-13-7P
     289038-15-9P
                    289038-16-0P
                                    289038-18-2P
                                                   289038-19-3P
                                                                   289038-21-7P
                    289038-24-0P
                                    289038-25-1P 289038-26-2P
     289038-23-9P
     289038-27-3P 289038-28-4P
                                  289038-29-5P
                                                 289038-30-8P
                    289038-33-1P
                                    289038-34-2P 289038-35-3P
     289038-32-0P
     289038-36-4P
     RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by
        cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as
        ileal bile acid transport and taurocholate uptake inhibitors)
IT
     289038-37-5P 289038-38-6P
                                  289038-39-7P
                                                 289038-40-0P
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289038-44-4P

289038-42-2P 289038-43-3P

289038-41-1P

289038-45-5P 289056-45-7P 289056-46-8P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as ileal bile acid transport and taurocholate uptake inhibitors)

197373-53-8P 280105-79-5P 280105-80-8P 280105-82-0P 280105-83-1P 280105-84-2P 280105-91-1P 280105-94-4P 280105-92-2P 280106-02-7P 280105-98-8P 280106-01-6P 280106-04-9P 280106-05-0P 280106-06-1P 280106-08-3P 280106-09-4P 280106-10-7P 280106-11-8P 280106-12-9P 289039-86-7P 289039-87-8P 289039-88-9P 289039-90-3P 289039-91-4P 289039-93-6P 289039-95-8P 289039-96-9P 289039-97-0P 289039-98-1P 289040-00-2P 289039-99-2P 289040-01-3P 647859-05-0P **647859-06-1P** 647859-03-8P 647859-04-9P 647859-07-2P 647859-08-3P 647859-09-4P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU

(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of substituted 5-aryl-benzothiepines by cyclization of

(preparation of substituted 5-aryl-benzothiepines by cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as ileal bile acid transport and taurocholate uptake inhibitors)

IT 197373-50-5P 197373-51-6P

RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as ileal bile acid transport and taurocholate uptake inhibitors)

RN 197373-50-5 HCAPLUS

IT

CM

1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-5-(3-nitrophenyl)-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197373-51-6 HCAPLUS

CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

IT 197373-37-8P 197373-54-9P 197374-04-2P 197374-59-7P 197375-96-5P 197376-42-4P 197376-55-9P 197384-36-4P 289038-26-2P 289038-27-3P 289038-28-4P 289038-35-3P 289038-36-4P 289038-37-5P 289038-38-6P 289038-43-3P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as ileal bile acid transport and taurocholate uptake inhibitors)

RN 197373-37-8 HCAPLUS

CN 1-Propanesulfonic acid, 3-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino], rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197373-54-9 HCAPLUS

CN 1-Pentanaminium, 5-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-5-oxo-, rel-, salt with trifluoroacetic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 197373-53-8 CMF C37 H60 N3 O4 S

Relative stereochemistry.

CM 2

CRN 14477-72-6 CMF C2 F3 O2

RN 197374-04-2 HCAPLUS
CN 2-Propenamide, N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

RN 197374-59-7 HCAPLUS

CN Propanamide, 3-bromo-N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197375-96-5 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-5-[2-(dimethylamino)phenyl]-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel-(9CI) (CA INDEX NAME)

RN 197376-42-4 HCAPLUS

CN Benzenaminium, 3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]-N,N,N-trimethyl-, iodide, rel-(9CI) (CA INDEX NAME)

Relative stereochemistry.

• I -

RN 197376-55-9 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-5-[3-(dimethylamino)phenyl]-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel-(9CI) (CA INDEX NAME)

RN 197384-36-4 HCAPLUS

CN 1-Hexanaminium, 6-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-6-oxo-, rel-, salt with trifluoroacetic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 197384-35-3 CMF C38 H62 N3 O4 S

Relative stereochemistry.

CM 2

CRN 14477-72-6 CMF C2 F3 O2

RN 289038-26-2 HCAPLUS

CN Pyridinium, 2-[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]methyl]-1-ethyl-, iodide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

• I-

RN 289038-27-3 HCAPLUS

CN Pyridinium, 3-[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]methyl]-1-ethyl-, iodide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

• I-

RN 289038-28-4 HCAPLUS

CN Pyridinium, 4-[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]methyl]-1-ethyl-, iodide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

• I

RN 289038-35-3 HCAPLUS

CN Pyridinium, 2-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-1-ethyl-, iodide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

• I-

RN 289038-36-4 HCAPLUS

CN Pyridinium, 3-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-1-ethyl-, iodide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

DI-

RN 289038-37-5 HCAPLUS

CN Benzenemethanaminium, 4-[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]sulfonyl]-N,N,N-triethyl-, iodide, rel- (9CI) (CA INDEX NAME)

• I-

RN 289038-38-6 HCAPLUS
CN Pyridinium, 4-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-1-ethyl-, iodide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

• I-

RN 289038-43-3 HCAPLUS
CN Benzenaminium, 3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]-N,N,N-triphenyl-, bromide, rel-(9CI) (CA INDEX NAME)

Relative stereochemistry.

● Br-

IT 197373-53-8P 280105-98-8P 647859-06-1P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of substituted 5-aryl-benzothiepines by cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as ileal bile acid transport and taurocholate uptake inhibitors)

RN 197373-53-8 HCAPLUS

CN 1-Pentanaminium, 5-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-5-oxo-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 280105-98-8 HCAPLUS

CN 1-Propanesulfonic acid, 3-[[3-[(3R,4S,5S)-3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 647859-06-1 HCAPLUS

CN 1-Pentanaminium, 5-[[3-[(3R,4S,5S)-3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-5-oxo-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

REFERENCE COUNT: . 231 THERE ARE 231 CITED REFERENCES AVAILABLE FOR

THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L9 ANSWER 5 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:376848 HCAPLUS

DOCUMENT NUMBER: 138:385315

TITLE: Mono- and di-fluorinated benzothiepines as inhibitors

of apical sodium co-dependent bile acid transport

(ASBT) and taurocholate uptake for treating

hyperlipidemic conditions and methods for preparation

INVENTOR(S): Koeller, Kevin J.; Tremont, Samuel J.

PATENT ASSIGNEE(S): G.D. Searle and Co., USA

SOURCE:

PCT Int. Appl., 589 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	PATENT NO.																				
WO	WO 2003040127					A1 20030515				WO 2002-US35257						20021104					
	W :	ΑE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,				
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,				
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KΕ,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,				
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,				
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TN,	TR,	TT,	TZ,				
		UA,	ŪĠ,	US,	UZ,	VC,	VN,	ΥU,	ZA,	ZM,	ZW,	AM,	AZ,	BY,	KG,	KZ,	MD,				
		RU,	TJ,	TM																	
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CA	2464	685			AA		2003	0515	1	CA 2	002-	2464		2	LU, MC, NL, GW, ML, MR, 20021104 20021104						
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US	6740	663			B2		2004	0525					•								
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EP	1448	546			A1		2004	0825		EP 2	002-	7787	11		2	0021	104				
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		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	SK						
JP	JP 2005518347						2005	0623	1	JP 2	003-	5421	73	20021104							
US	2004	1764	38		A1		2004	0909	,	US 2	003-	7434	04		2	0031	223				
PRIORIT	PRIORITY APPLN. INFO.:								,	US 2	001-	3308	92P		P 2	0011	102				
										US 2	002-	2869	87		A3 2	0021	104				
						1	WO 2	002-	US35:	257		W 2	0021	104							
OTHER C	OTTROE		MAD.	ידיעם	138.	2052	1 5														

OTHER SOURCE(S):

MARPAT 138:385315

Ι

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AB Mono-fluorinated and di-fluorinated benzothiepine apical Na co-dependent bile acid transport (ASBT) inhibitors (shown as I; variables defined below; no specific examples are included) are disclosed together with methods of making the same, methods of using the same to treat hyperlipidemic conditions as well as pharmaceutical compns. containing the same compds. For I: X = F, X' = H, F; n = 0-2; m = 0-4; R2A and R2B = Hand hydrocarbyl; R3A, R3B, R5A, and R5B = H, alkyl, cycloalkyl, alkenyl, alkynyl, heterocyclyl, quaternary heterocyclyl, oxo, aryl-R5, -OR9, -NR9R10, -SR9, -S(0)R9, -SO2R9, and -SO3R9; R9 and R10 = H, hydrocarbyl, amino, and hydrocarbylamino. R5 = H, hydrocarbyl, heterocyclyl,

quaternary heterocyclyl, -OR9, -SR9, -S(O)R9, -SO2R9, and -SO3R9; ≥1 R6 radicals = H, halogen, -CN, -NO2, hydrocarbyl, -R5, -OR13, -NR 13R14, -SR13, -S(0)R13, -S(0)2R13, -S03R13, -S+R3R14A-, -NR13OR14, -NR13NR14R15, -OM, -SO2OM, -SO2NR13R14, -NR14C(O)R13, -C(O)OM, -S(O)NR13R14, -N+R13R14R15A-, -PR13R14, -P(O)R13R4, -P+R13R14R15A-, amino acid residue, peptide residue, polypeptide residue, and carbohydrate residue; addnl. details are given in the claims. I (X = X' = F) are claimed to be preparable from the 4-oxo analog and diethylaminosulfur trifluoride; I (X = F; X' = H) are claimed preparable from the 4-hydroxy analog and diethylaminosulfur trifluoride. Hundreds of example prepns. of precursors to I are included, but none of I; most of the example prepns. have appeared in earlier patents (e.g. WO 98/40375). Biol. testing procedures are described but no test results are reported except for the statement that a polyethylene glycol-functionalized benzothiepine (4500 MW; a 4-hydroxy analog of I) inhibited ileal bile acid transport-mediated uptake of 14C-taurocholate in H14 cells.

- IC ICM C07D337-00 ICS A61K031-38
- CC 27-21 (Heterocyclic Compounds (One Hetero Atom)) Section cross-reference(s): 1

289037-96-3P 289037-98-5P

1426-54-6P, 4-Fluoro-2-[(4-methoxyphenyl)methyl]phenol IT 1481-12-5P, 4-Fluoro-2-(4'-fluorobenzyl)phenol 1515-89-5P, 3-Bromobenzyl methyl 3670-91-5P 15886-84-7P 16473-35-1P, 1-(Chloromethyl)-4-(hydroxymethyl)benzene 24632-01-7P, 1-(Hydroxymethyl)cyclohexanecarboxal 24765-57-9P, 2,2-Dibutyl-1,3-propanediol 70132-87-5P dehyde 120454-34-4P, 2-Mercaptodiphenylmethane 120936-00-7P, O-2-Benzylphenyl 120936-01-8P 131117-88-9P 162632-54-4P, dimethylthiocarbamate 2-Mercapto-4-methoxybenzophenone 163445-43-0P, 2-Mercapto-5methoxybenzophenone 174747-95-6P, 1-Bromo-2-butyl-2-(hydroxymethyl)hexane 178678-21-2P 178678-22-3P, 3-Butyl-3-ethyl-5phenyl-2,3-dihydrobenzothiepine 178678-23-4P, cis-3-Butyl-3-ethyl-5phenyl-2,3-dihydrobenzothiepin-4(5H)-one 178678-24-5P, trans-3-Butyl-3-ethyl-5-phenyl-2,3-dihydrobenzothiepin-4(5H)-one 178678-25-6P, cis-3-Butyl-3-ethyl-5-phenyl-2,3-dihydrobenzothiepin-4(5H)one-1,1-dioxide 178678-26-7P 178678-27-8P 178678-29-0P 178678-33-6P, 3-Ethyl-5-phenyl-2,3-dihydrobenzothiepine 178678-34-7P 178678-36-9P, cis-3-Ethyl-5-phenyl-2,3,4,5-tetrahydrobenzothiepine-1,1-178678-37-0P 178678-40-5P 178678-45-0P 178678-46-1P dioxide 178678-50-7P 178678-51-8P 178678-55-2P 178678-56-3P, 178678-49-4P 2-[(2-Benzoylphenylthio)methyl]-2-ethylhexanal 178678-57-4P, 2-[(2-Benzoylphenylthio)methyl]butyraldehyde 178678-58-5P 178678-59-6P 178678-60-9P 178678-61-0P 178678-62-1P 178678-63-2P 178678-64-3P 178678-65-4P 178678-66-5P 178678-67-6P 178678-68-7P 178678-69-8P 178678-70-1P 178678-71-2P 178678-72-3P 178678-73-4P 178897-97-7P 178898-01-6P 178898-05-0P 197372-67-1P 197372-71-7P 178898-00-5P 197372-76-2P 197372-77-3P 197372-78-4P 197373-02-7P 197373-03-8P 197373-13-0P 197373-42-5P 197373-43-6P 197373-44-7P 197373-46-9P 197373-47-0P 197373-49-2P 197373-50-5P 197373-51-6P 197373-52-7P 197373-57-2P 197373~55-0P 197373-56-1P 197373-58-3P 197377-84-7P 197378-05-5P 197378-07-7P, 4-Chloro-3-(4-methoxyphenylmethyl)nitrobenzene 197378-15-7P 197378-16-8P 197378-18-0P 197378-20-4P 197378-22-6P 197378-24-8P 197378-26-0P 197378-29-3P 197378-31-7P 197378-32-8P 197378-34-0P 197378-36-2P 197378-38-4P 197378-40-8P 197378-42-0P 197378-44-2P 197378-46-4P 197378-48-6P, 4-Fluoro-2-(3'-methoxybenzyl)phenol 197378-50-0P 197378-52-2P 197378-54-4P 197378-56-6P 197378-58-8P 228113-64-2P 228113-57-3P 228113-58-4P 228113-59-5P 228113-63-1P 270931-13-0P 270931-14-1P 270931-15-2P 288863-77-4P

289038-47-7P

289038-46-6P

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                                                                 289038-84-2P
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                   525589-61-1P
                                  525589-62-2P
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                                                                 525589-64-4P
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     525589-71-3P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation of precursors of mono- and di-fluorinated benzothiepine
        inhibitors of apical sodium co-dependent bile acid transport (ASBT) and
        taurocholate uptake for treating hyperlipidemic conditions)
     178678-28-9P, 3-Butyl-3-ethyl-5-phenyl-2,3-dihydrobenzothiepine-1,1-
IT
               178678-30-3P, cis-3-Butyl-3-ethyl-5-phenyl-2,3,4,5-
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     tetrahydrobenzothiepine-1,1-dioxide
                                           178678-31-4P, trans-3-Butyl-3-ethyl-
     5-phenyl-2,3,4,5-tetrahydrobenzothiepine-1,1-dioxide
                                                           178678-32-5P,
     3-Butyl-3-ethyl-4-hydroxy-5-cyclohexylidene-2,3,4,5-
     tetrahydrobenzothiepine-1,1-dioxide
                                           178678-35-8P
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                   178678-41-6P
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                                                  178898-04-9P
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     525589-59-7P
                   526199-85-9P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of precursors of mono- and di-fluorinated benzothiepine
        inhibitors of apical sodium co-dependent bile acid transport (ASBT) and
        taurocholate uptake for treating hyperlipidemic conditions)
IT
     197373-50-5P 197373-51-6P 197373-52-7P
     289037-96-3P 289037-98-5P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation of precursors of mono- and di-fluorinated benzothiepine
        inhibitors of apical sodium co-dependent bile acid transport (ASBT) and
        taurocholate uptake for treating hyperlipidemic conditions)
RN
     197373-50-5 HCAPLUS
CN
     1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-5-(3-
     nitrophenyl)-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)
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RN 197373-51-6 HCAPLUS
CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3,3-dibutyl-7-(dimethylamino)2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197373-52-7 HCAPLUS
CN Pentanamide, 5-bromo-N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

RN 289037-96-3 HCAPLUS

CN Carbamic acid, [3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, 3-chloropropyl ester, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 289037-98-5 HCAPLUS

CN Urea, N-[3-(chloromethyl)phenyl]-N'-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

IT 197373-54-9P 280105-90-0P 289037-97-4P 289037-99-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of precursors of mono- and di-fluorinated benzothiepine inhibitors of apical sodium co-dependent bile acid transport (ASBT) and taurocholate uptake for treating hyperlipidemic conditions)

RN 197373-54-9 HCAPLUS

CN 1-Pentanaminium, 5-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-5-oxo-, rel-, salt with trifluoroacetic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 197373-53-8 CMF C37 H60 N3 O4 S

Relative stereochemistry.

CM 2

CRN 14477-72-6 CMF C2 F3 O2

RN 280105-90-0 HCAPLUS

CN Poly(oxy-1,2-ethanediyl),  $\alpha$ -[2-[[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]carbonyl]amino]ethyl]- $\omega$ -methoxy-, rel- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{NMe}_2 \\ \text{MeO} \\ \hline \end{array} \\ \begin{array}{c} \text{CH}_2 - \text{CH}_2 - \text{O} \\ \hline \end{array} \\ \begin{array}{c} \text{N} \\ \text{CH}_2 - \text{CH}_2 - \text{NH} \\ \hline \end{array} \\ \begin{array}{c} \text{CH}_2 - \text{CH}_2 - \text{NH} \\ \hline \end{array} \\ \begin{array}{c} \text{N} \\ \text{O} \end{array}$$

RN 289037-97-4 HCAPLUS

CN 4-Aza-1-azoniabicyclo[2.2.2]octane, 1-[3-[[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]carbonyl]oxy]propyl]-, chloride, rel- (9CI) (CA INDEX NAME)

$$n-Bu$$
 $R$ 
 $R$ 
 $NMe_2$ 
 $HO$ 
 $CCH_2)_3$ 

• c1-

RN 289037-99-6 HCAPLUS

CN 4-Aza-1-azoniabicyclo[2.2.2]octane, 1-[[3-[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]carbonyl]amino]phenyl]methyl]-, chloride, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

● Cl -

REFERENCE COUNT:

5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 6 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: DOCUMENT NUMBER:

2003:173440 HCAPLUS

138:215326

TITLE:

Combined preparations, containing 1,4-benzothiepine-1,1-dioxide derivatives and other active substances

for the treatment of hyperlipidemia

INVENTOR(S):

Glombik, Heiner; Frick, Wendelin; Schaefer,

Hans-Ludwig; Kramer, Werner

PATENT ASSIGNEE(S):

Aventis Pharma Deutschland GmbH, Germany

SOURCE:

PCT Int. Appl., 40 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

GI

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

			KIND DATE								DATE						
	WO 2003018024										2002-		20020809				
											, BG,					CH,	CN,
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC	, EE,	ES,	FI,	GB,	GD,	GE,	GH,
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		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN	, MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
		PL,	PT,	RO,	RŪ,	SD,	SE,	SG,	SI,	SK	, SL,	TJ,	TM,	TN,	TR,	TT,	TZ,
		UA,	ŪĠ,	UΖ,	VN,	YU,	ZA,	ZM,	ZW								
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DE	DE 10140169					A1 20030306				DE	2001-	1014	20010822				
	DE 10142456																
CA	2457	AA 20030306				CA	2002-	2457									
EP	1425018											20020809					
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	BR 2002012031						A 20040803				2002-	1203	20020809				
JP	JP 2005501861						T2 20050120				2003-	5225	20020809				
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	NO 2004000702						A 20040519										
PRIORIT	PRIORITY APPLN. INFO.:										2001-						
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OTHER S	MAR	MARPAT 138:21532															

$$R^4$$
 $R^5$ 
 $R^2$ 
 $R^2$ 
 $R^3$ 

Ι

- The invention relates to mixts. of substances, containing 1,4-benzothiepine1,1-dioxide derivs. of formula (I), in which the functional groups have
  the indicated meanings, their physiol. acceptable salts and physiol.
  functional derivs. as well as other active substances for the treatment of
  metabolic disorders especially hyperlipidemia. The combinations can also
  include antidiabetics, antiarthrytics etc. A typical capsule contains 100
  mg of the drugs and 400 mg triglyceride mixture from coco fatty acids; other
  formulations are emulsions, tablets, dragees, and solns. Hamster that
  were fed with cholesterol-rich feed received orally the drug combination
  once daily for 10 days. Feces was analyzed for bile acids, blood lipid
  levels were measured and cholesterol was determined from liver.
- IC ICM A61K031-55

ICS A61K031-395; A61P003-06

CC 1-10 (Pharmacology)

Section cross-reference(s): 63

IT 56-03-1, Biguanide 300-62-9, Amphetamine 943-45-3, Fibric acid 2295-31-0, Glitazone 5395-30-2 9000-40-2, Carob gum 9002-72-6, Growth hormone 9004-10-8, Insulin, biological studies 9034-39-3, Growth hormone releasing hormone 11041-12-6, Cholestyramine 25614-03-3, Bromocriptine 25812-30-0, Gemfibrozil 49642-07-1, Statine 50925-79-6, Cholestipol 54870-28-9, Meglitinide 96829-58-2, Orlistat 99759-19-0, Tiqueside 129024-87-9, Doprexin 150332-35-7, Pamaqueside 163222-33-1, Ezetimibe 252047-40-8

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(combined prepns., containing 1,4-benzothiepine-1,1-dioxide derivs. and other active substances for treatment of hyperlipidemia)

IT 252047-40-8

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(combined prepns., containing 1,4-benzothiepine-1,1-dioxide derivs. and other active substances for treatment of hyperlipidemia)

RN 252047-40-8 HCAPLUS

CN D-Glucitol, 1-[[5-[[3-[(3S,4R,5R)-3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-5-oxopentyl]amino]-1-deoxy-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

$$R = R$$

NMe<sub>2</sub>

HN

(CH<sub>2</sub>) 4

NMe<sub>2</sub>

OH

OH

OH

OH

OH

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 7 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:487559 HCAPLUS

DOCUMENT NUMBER: 137:63115

TITLE: Preparation of diphenylazetidinone derivatives as

hypolipidemic agents

INVENTOR(S): Glombik, Heiner; Kramer, Werner; Flohr, Stefanie;

Frick, Wendelin; Heuer, Hubert; Jaehne, Gerhard;

Lindenschmidt, Andreas; Schaefer, Hans-Ludwig

PATENT ASSIGNEE(S): Aventis Pharma Deutschland GmbH, Germany

SOURCE: PCT Int. Appl., 67 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

										APPLICATION NO.											
	WO					A1 20020627								20011211							
		W:	ΑE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BE	3,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,		
			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC	Ξ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,		
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE	Ξ,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,		
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN	1,	MW,	MX,	MZ.	NO,	NZ,	OM,	PH,		
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						YU,				•		•	•	·	•	•	•	•	•		
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	DE	10064	•																		
						Δ1	DE 2000-10064402 DE 2001-10154520							20001221							
	CA	2431	985							CA 2001-10134320											
										AU 2001-2431983											
	DD	2002	0033.	73		A 20020701				EE 2002-131/3							20011211				
	EE	1245	222	,		A 20030013			EE 2003-237 EP 2001-271371							20011211					
	EP							GB, GR, IT, LI, LU,													
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	JP	P 2004516293					7 20041126			NZ 2001-526592							20011211				
	147	NZ 320332																			
							RU 2003-122219														
	US 2002128252										US 2001-21028						2	0011	219		
	US 6498156							2002											_		
	ZA 2003004092							2004			ZA	20	03-4	4092			2	0030			
	ZA 2003004095 .										ZA	20	03-4	4095			2	0030			
	NO 2003002733										NO	20	03-2	2733			2	0030	616		
	HK 1059936							2006	0127		HK	20	04-	1028	49		2	0040			
PRIOR	PRIORITY APPLN. INFO.:																	0001	221		
											DΕ	20	01-	1015	4520		A 2	0011	107		
											WO	20	01-1	EP14	532		W 2	0011	211		
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OTHER SOURCE(S): MARPAT 137:63115

GΙ

<sup>\*</sup> STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

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The compds. are suited for use e.g. as hypolipidemic drugs. The invention
AB
     discloses preparation of diphenylazetidinone derivs. such as I [R1, R2, R3, R4,
     R5, R6 = C0-C30-alkylene-L {optionally containing O, CO, CH:CH, C.tplbond.C,
     N(alkyl), N(alkylphenyl), NH}, H, F, Cl, Br, I, CF3, NO2, CN, CO2H,
     CO2(alkyl), CONH2, CONH(alkyl), CON(alkyl)2, alkyl, alkenyl, alkynyl,
     O-alkyl, SO2NH2, SO2NH(alkyl) SO2N(alkyl)2, S-(alkyl), SO(alkyl),
     (un) substituted S(CH2) nPh, SO(CH2) nPh, SO2(alky1), SO2(CH2) nPh, NH2,
     NH(alkyl), N(alkyl)2, NH(acyl), (un) substituted Ph, O(CH2) nPh; n = 0-6; L
     = II; R7, R9, R10 = Me, Et, Pr, butyl; R8 = H, OH, NH2, NH(alkyl)], and
     their physiol. acceptable salts, for their use as hypolipidemic agents.
     Thus, 1,2-diphenylazetidinone derivative III trifluoroacetate (IV) was
     prepared via a multistep synthetic sequence starting from
     7-[3-(3-butyl-7-dimethylamino-3-ethyl-4-hydroxy-1,1-dioxo-2,3,4,5-
     tetrahydro-1H-benzo[b]thiepin-5-yl)-phenylcarbamoyl]-heptanoic acid and
     4-(4-aminomethylphenyl)-1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-
     hydroxyphenyl]-azetidin-2-one. Azetidinone IV was tested for its
     cholesterol lowering ability [ED50 = 0.01 mg/mouse].
IC
     ICM C07D409-12
     ICS A61K031-397; A61P009-00
     26-5 (Biomolecules and Their Synthetic Analogs)
CC
     Section cross-reference(s): 1, 63
IT
     439113-82-3P 439113-89-0P 439113-91-4P
     439113-92-5P 439113-93-6P 439113-96-9P
     439113-98-1P 439114-01-9P 439114-03-1P
     439114-06-4P 439114-08-6P 439114-11-1P
     439114-16-6P 439114-20-2P 439114-22-4P
     439114-23-5P 439114-26-8P 439114-29-1P
     439114-36-0P 439114-38-2P 439114-39-3P
     439114-40-6P 439120-25-9P
     RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (preparation of diphenylazetidinone derivs. as hypolipidemics)
     76-05-1, Trifluoroacetic acid, reactions 112-60-7, Tetraethylene glycol
IT
     124-04-9, Hexanedioic acid, reactions 1117-97-1, O,N-
     Dimethylhydroxylamine 1501-05-9 1663-39-4, tert-Butyl acrylate
     7480-32-2, 4-Phenyl-oxazolidin-2-one 20256-89-7 23243-68-7
     402820-38-6
                 439080-96-3 439114-09-7 439114-17-7
     439114-41-7 439114-42-8 439114-43-9
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (preparation of diphenylazetidinone derivs. as hypolipidemics)
IT
     94376-75-7P 439080-20-3P 439080-21-4P
                                                439080-24-7P 439080-59-8P
     439080-60-1P
                   439080-61-2P
                                  439080-62-3P
                                                  439113-83-4P
                                                                 439113-84-5P
     439113-85-6P
                   439113-86-7P
                                  439113-87-8P 439113-88-9P
     439113-90-3P 439113-94-7P 439113-99-2P
                                  439114-13-3P 439114-14-4P
     439114-04-2P
                  439114-12-2P
     439114-18-8P 439114-24-6P 439114-27-9P
                  439114-31-5P 439114-32-6P 439114-34-8P
     439114-30-4P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation of diphenylazetidinone derivs. as hypolipidemics)
IT
     439113-82-3P 439113-89-0P 439113-91-4P
     439113-92-5P 439113-93-6P 439113-96-9P
     439113-98-1P 439114-01-9P 439114-03-1P
     439114-06-4P 439114-08-6P 439114-11-1P
     439114-16-6P 439114-20-2P 439114-22-4P
     439114-23-5P 439114-26-8P 439114-29-1P
     439114-36-0P 439114-38-2P 439114-39-3P
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## 439114-40-6P 439120-25-9P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of diphenylazetidinone derivs. as hypolipidemics) 439113-82-3 HCAPLUS

RN 439113-82-3 HCAPLUS
CN Pentanamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-5-[[[4-[3-(3-hydroxy-3-phenylpropyl)-2-(4-methoxyphenyl)-4-oxo-1-azetidinyl]phenyl]methyl]amino]-(9CI) (CA INDEX NAME)

PAGE 1-B

− NMe<sub>2</sub>

RN 439113-89-0 HCAPLUS

CN Hexanediamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-N'-[[4-[1-(4-fluorophenyl)-3-(3-hydroxy-3-phenylpropyl)-4-oxo-2-azetidinyl]phenyl]methyl]- (9CI) (CA INDEX NAME)

RN 439113-91-4 HCAPLUS

CN Hexanediamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-N'-[[4-[1-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]- (9CI) (CA INDEX NAME)

PAGE 1-A

RN 439113-92-5 HCAPLUS

CN Hexanediamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-N'-[[4-[3-(3-hydroxy-3-phenylpropyl)-2-(4-methoxyphenyl)-4-oxo-1-azetidinyl]phenyl]methyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

RN 439113-93-6 HCAPLUS

CN 5,8,11-Trioxa-2-azatridecan-13-amide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-1-[4-[1-(4-fluorophenyl)-3-(3-hydroxy-3-phenylpropyl)-4-oxo-2-azetidinyl]phenyl]-3-oxo- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

RN 439113-96-9 HCAPLUS

CN Acetamide, 2-[2-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-2-oxoethoxy]ethoxy]-N-[[4-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]-, mono(trifluoroacetate)(salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439113-95-8

CMF C55 H64 F2 N4 O9 S

PAGE 1-A

PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439113-98-1 HCAPLUS

CN Acetamide, 2-[2-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-2-oxoethoxy]ethoxy]-N-[[3-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]-, mono(trifluoroacetate)

(salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439113-97-0 CMF C55 H64 F2 N4 O9 S

PAGE 1-A

PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-01-9 HCAPLUS

CN 5,8,11-Trioxa-2-azatridecan-13-amide, N-[3-[3-butyl-7-(dimethylamino)-3-

ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-1-[4-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]-3-oxo-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-00-8

CMF C57 H68 F2 N4 O10 S

PAGE 1-A

PAGE 1-B

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-03-1 HCAPLUS

CN 5,8,11-Trioxa-2-azatridecan-13-amide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-1-[3-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]-3-oxo-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-02-0 CMF C57 H68 F2 N4 O10 S

PAGE 1-A

PAGE 1-B

CM 2

CRN 76-05-1

CMF C2 H F3 O2

RN 439114-06-4 HCAPLUS

CN Dodecanediamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-N'-[[4-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-05-3 CMF C61 H76 F2 N4 O7 S

PAGE 1-A

$$\begin{array}{c} & & \text{Et} \\ & & \text{n-Bu} \\ & & \text{N} \\ & & \text{HO} \\ & & \text{CH}_2 - \text{NH} - \text{C} - (\text{CH}_2)_{10} - \text{C} - \text{NH} \\ & & \text{O} \\ & & & \text{O} \\ &$$

PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-08-6 HCAPLUS

CN Dodecanediamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-N'-[[3-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-07-5 CMF C61 H76 F2 N4 O7 S

PAGE 1-A

PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-11-1 HCAPLUS

CN 5,8,11-Trioxa-2-azatridecan-13-amide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-1-[4-[3-(3-hydroxy-3-phenylpropyl)-2-(4-methoxyphenyl)-4-oxo-1-azetidinyl]phenyl]-3-oxo-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-10-0 CMF C58 H72 N4 O11 S

PAGE 1-A

PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-16-6 HCAPLUS

CN 4,7,10,13,16-Pentaoxanonadecanediamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-N'-[[3-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-15-5 CMF C63 H80 F2 N4 O12 S

PAGE 1-A

PAGE 1-B

— 
$$\mathrm{CH_2}-\mathrm{O}-\mathrm{CH_2}-\mathrm{CH_2}-\mathrm{O}-\mathrm{CH_2}-\mathrm{CH_2}-\mathrm{O}-\mathrm{CH_2}-\mathrm{$$

PAGE 1-C

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-20-2 HCAPLUS

CN 4,7,10,13,16,19,22-Heptaoxapentacosanediamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-N'-[[3-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-19-9 CMF C67 H88 F2 N4 O14 S Krishnan 10/734 '87

09/05/2006

PAGE 1-A

PAGE 1-B

$$-$$
 CH2 $-$  O $-$  CH2 $-$  $-$  CH2

PAGE 1-C

$$\begin{array}{c|c} & & & & \\ & &$$

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-22-4 HCAPLUS

CN 4,7,10,13,16-Pentaoxanonadecanediamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-N'-[[4-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-21-3.

CMF C63 H80 F2 N4 O12 S

PAGE 1-A

PAGE 1-B

PAGE 1-C

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-23-5 HCAPLUS

CN Octanoic acid, 8-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-8-oxo-(9CI) (CA INDEX NAME)

RN 439114-26-8 HCAPLUS

CN Octanamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-8-[[4-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]amino]-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-25-7 CMF C57 H70 F2 N4 O6 S

$$\begin{array}{c} \text{Et} \\ \text{n-Bu} \\ \text{NMe}_2 \\ \text{HO} \\ \text{CH}_2-\text{NH}-\text{(CH}_2)_{\,7}-\text{C-NH} \\ \text{O} \\ \text{OH} \\ \text{F} \end{array}$$

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-29-1 HCAPLUS

CN Acetamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-2-[2-[2-[[4-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]amino]ethoxy]ethoxy]-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-28-0 CMF C55 H66 F2 N4 O8 S

PAGE 1-A

PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-36-0 HCAPLUS
CN Acetamide, 2-[2-[2-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]ethoxy]ethoxy]-N-[[4-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]-,

mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM I

CRN 439114-35-9

CMF C55 H66 F2 N4 O8 S

PAGE 1-A

PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-38-2 HCAPLUS

CN Acetamide, 2-[2-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]ethoxy]ethoxy]-N-[[3-[1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-37-1 CMF C55 H66 F2 N4 O8 S

PAGE 1-A

PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 439114-39-3 HCAPLUS

CN Acetamide, 2-[2-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-2-oxoethoxy]ethoxy]-N-[[4-[3-[3-(4-fluorophenyl)-3-hydroxypropyl]-2-(4-methoxyphenyl)-4-oxo-1-azetidinyl]phenyl]methyl]- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

RN 439114-40-6 HCAPLUS

CN 5,8,11-Trioxa-2-azatridecan-13-amide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-1-[4-[3-[3-(4-fluorophenyl)-3-hydroxypropyl]-2-(4-methoxyphenyl)-4-oxo-1-azetidinyl]phenyl]-3-oxo- (9CI) (CA INDEX NAME)

PAGE 1-A

OMe
$$CH_2-NH-C-CH_2-O-CH_2-CH_2-$$

$$CH-CH_2-CH_2$$

PAGE 1-B

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 439120-25-9 HCAPLUS

CN Acetamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-2-[2-[2-[[[3-[1-(4-fluorophenyl)-3-hydroxypropyl]-4-oxo-2-azetidinyl]phenyl]methyl]amino]ethoxy]ethoxy]-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439120-24-8

CMF C55 H66 F2 N4 O8 S

## PAGE 1-A

$$\begin{array}{c} \text{Et} \\ \text{n-Bu} \\ \text{HO} \\ \\ \text{CH}_2-\text{NH}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{C}-\text{NH}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{C}-\text{NH}-\text{CH}_2-\text{CH}_2-\text{CH}_2-\text{C}-\text{NH}-\text{CH}_2-\text{CH}$$

## PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

CN

IT 439114-09-7 439114-42-8 439114-43-9

RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of diphenylazetidinone derivs. as hypolipidemics)

RN 439114-09-7 HCAPLUS

Undecanediamide, N'-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-N-methoxy-N-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & \\ & & \\$$

RN 439114-42-8 HCAPLUS
CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3-butyl-7-(dimethylamino)-3-ethyl2,3,4,5-tetrahydro-, 1,1-dioxide (9CI) (CA INDEX NAME)

RN 439114-43-9 HCAPLUS
CN Pentanamide, 5-bromo-N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]- (9CI) (CA INDEX NAME)

IT 439113-88-9P 439113-94-7P 439113-99-2P 439114-04-2P 439114-14-4P 439114-18-8P 439114-24-6P 439114-27-9P 439114-32-6P 439114-34-8P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of diphenylazetidinone derivs. as hypolipidemics)

RN 439113-88-9 HCAPLUS

CN Hexanoic acid, 6-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-6-oxo-(9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{DO} \\ \text{DO} \\ \text{NMe}_2 \\ \text{HO}_2\text{C-} \\ \text{CH}_2 \\ \text{O} \end{array}$$

RN 439113-94-7 HCAPLUS

CN Acetic acid, [2-[2-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-2-oxoethoxy]ethoxy]- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{D} \\ \text{NMe}_2 \\ \text{HO}_2\text{C--} \text{CH}_2\text{--} \text{O--} \text{CH}_2\text{--} \text{C--} \text{NH} \\ \\ \text{O} \\ \end{array}$$

RN 439113-99-2 HCAPLUS

CN Acetic acid, [2-[2-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-2-oxoethoxy]ethoxy]- (9CI) (CA INDEX NAME)

$$\begin{array}{c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

RN 439114-04-2 HCAPLUS

CN Dodecanoic acid, 12-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-12-oxo-(9CI) (CA INDEX NAME)

RN 439114-14-4 HCAPLUS

CN 4,7,10,13,16-Pentaoxanonadecanoic acid, 19-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-19-oxo- (9CI) (CA INDEX NAME)

PAGE 1-A

 $\text{HO}_2\text{C}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH}_2-\text{CH}_2-\text{O}-\text{CH}_2-\text{CH$ 

PAGE 1-B

$$\begin{array}{c|c} & & & & \\ & &$$

RN 439114-18-8 HCAPLUS

CN 4,7,10,13,16,19,22-Heptaoxapentacosanoic acid, 25-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-25-oxo-(9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

RN 439114-24-6 HCAPLUS

CN Octanediamide, N'-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-N-methoxy-N-methyl-(9CI) (CA INDEX NAME)

RN 439114-27-9 HCAPLUS

CN 2,6,9-Trioxa-3-azaundecan-11-amide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-3-methyl-4-oxo-(9CI) (CA INDEX NAME)

$$\begin{array}{c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

RN 439114-32-6 HCAPLUS

CN Acetic acid, [2-[2-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]ethoxy]ethoxy]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

RN 439114-34-8 HCAPLUS

CN Acetic acid, [2-[2-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]ethoxy]ethoxy]-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 439114-33-7 CMF C30 H44 N2 O7 S

CM 2

CRN 76-05-1 CMF C2 H F3 O2

REFERENCE COUNT:

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 8 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

5

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ACCESSION NUMBER: 2001:693092 HCAPLUS

DOCUMENT NUMBER: 135:257253

TITLE: Preparation of tetrahydrobenzothiepines and

naphthalenes useful in combination therapy
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naphthalenes useful in combination therapy with HMG Co-A reductase inhibitors for the prophylaxis and treatment of hyperlipidemic conditions and disorders. Keller, Bradley T.; Tremont, Samuel J.; Glenn, Kevin

C.; Manning, Robert E.
PATENT ASSIGNEE(S): Pharmacia Corporation, USA
SOURCE: PCT Int. Appl., 182 pp.

.:.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

INVENTOR (S):

PA'	PATENT NO.							APPLICATION NO.			DATE						
				A2 20010920			WO 2001-US7505			20010308							
	W:	AE, CR, HU, LU, SD, YU,	AG, CU, ID, LV, SE, ZA,	AL, CZ, IL, MA, SG, ZW,	AM, DE, IN, MD, SI, AM,	AT, DK, IS, MG, SK, AZ,	DM, JP, MK, SL, BY,	DZ, KE, MN, TJ, KG,	EE, KG, MW, TM, KZ,	ES KP MX TR MD	, BG, , FI, , KR, , MZ, , TT, , RU, , TZ,	GB, KZ, NO, TZ, TJ,	GD, LC, NZ, UA, TM	GE, LK, PL, UG,	GH, LR, PT, US,	GM, LS, RO, UZ,	HR, LT, RU, VN,
US	2002	BJ, 0618 2328	CF, 88 34	CG,	CI, A1 A1	CM,	GA, 2002 2003	GN, 0523 1218	GW, t	ML JS JS	, LU, , MR, 2001-8	NE, 3023: 2046	SN, 13 72	TD,	TG 2	0010 0021	308 126
	2004 2004 Y APP	1107	61						; ; ; ;	US US US US	2003-( 2003-( 2000-) 2000-) 2001-( 2001-)	51194 1883 1883 3022 3023	42 61P 78P 79	] ] ]	2 P 2 P 2 A3 2 B1 2	0030 0030 0000 0000 0010 0010	703 310 310 308 308

- AB A method for the treatment and/or prophylaxis of a hyperlipidemic condition or disorder comprises the administration of ≥1 HMG Co-A reductase inhibitors and one or more specific apical Na codependent bile acid transporter (ASBT) inhibitors is claimed. Thus, (4R,5R)-1-[[4-[4-[3-butyl-3-ethyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxo-1-benzothiepin-5-yl]phenoxy]methyl]phenyl]methyl-4-aza-1-azoniabicyclo[2.2.2]octane chloride (3,3-di-Bu analog preparation given) 0.375 mg/kg/day and lovastatin 0.45 mg/kg/day orally in dogs reduced serum triglycerides by 37% at 4 wk.
- IC ICM A61K031-495
  - ICS A61K031-38; A61K031-235
- CC 28-16 (Heterocyclic Compounds (More Than One Hetero Atom))
  Section cross-reference(s): 1, 25, 27, 63
- 73573-88-3, Mevastatin 75330-75-5, Lovastatin 79902-63-9, Simvastatin TΤ 81093-37-0, Pravastatin 93957-54-1, Fluvastatin 134523-00-5, Atorvastatin 145599-86-6, Cerivastatin 147098-20-2, ZD-4522 147526-32-7 151165-96-7 280105-82-0 280105-83-1 280105-84-2 **280105-88-6 280105-89-7 280105-91-1 280105-92-2** 280757-38-2 289037-90-7 289039-91-4 361484-19-7 361484-23-3 361484-26-6

RL: BAC (Biological activity or effector, except adverse); BSU (Biological

study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(preparation of tetrahydrobenzothiepines and naphthalenes useful in combination therapy with HMG Co-A reductase inhibitors for the prophylaxis and treatment of hyperlipidemic conditions and disorders)

25784-91-2P, 2-Chloro-5-nitrobenzoyl chloride 16473-35-1P 24765-57-9P 70132-87-5P 174747-95-6P 197373-49-2P 197373-50-5P 197373-51-6P 197373-55-0P 197373-56-1P 197373-57-2P 197378-07-7P 197378-31-7P 197378-32-8P 197378-46-4P 197378-48-6P 197378-52-2P 197378-50-0P 197378-54-4P 197378-56-6P 197378-58-8P 228113-65-3P 361373-66-2P 361373-79-7P 361373-81-1P 361373-83-3P 361373-85-5P 361373-87-7P 361373-89-9P 361373-91-3P 361373-92-4P 361373-94-6P 361373-96-8P 361373-98-0P 361374-00-7P 361374-02-9P

361484-28-8P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of tetrahydrobenzothiepines and naphthalenes useful in combination therapy with HMG Co-A reductase inhibitors for the prophylaxis and treatment of hyperlipidemic conditions and disorders) 280105-88-6 280105-89-7

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES

(preparation of tetrahydrobenzothiepines and naphthalenes useful in combination therapy with HMG Co-A reductase inhibitors for the prophylaxis and treatment of hyperlipidemic conditions and disorders) 280105-88-6 HCAPLUS

Pyridinium, 4-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-1methyl-, rel-, salt with 4-methylbenzenesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

IT

IT

RNCN

> CRN 280105-87-5 CMF C32 H44 N3 O3 S

Relative stereochemistry.

361374-06-3P

361374-08-5P

CM 2

CRN 16722-51-3 CMF C7 H7 O3 S

RN 280105-89-7 HCAPLUS

CN 4-Aza-1-azoniabicyclo[2.2.2]octane, 1-[[4-[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]methyl]phenyl]methyl]-, chloride, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

• c1 ~

IT 197373-50-5P 197373-51-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of tetrahydrobenzothiepines and naphthalenes useful in combination therapy with HMG Co-A reductase inhibitors for the prophylaxis and treatment of hyperlipidemic conditions and disorders) 197373-50-5 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-5-(3-nitrophenyl)-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN

RN 197373-51-6 HCAPLUS

CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

L9 ANSWER 9 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:560070 HCAPLUS

DOCUMENT NUMBER: 135:137410

TITLE: Preparation of ileal bile acid transport inhibiting

benzothiepines for combination therapy with HMG Co-A

reductase inhibitors.

INVENTOR(S): Keller, Bradley T.; Glenn, Kevin C.; Manning, Robert

Ε.

PATENT ASSIGNEE(S): G.D. Searle and Co., USA

SOURCE: U.S., 356 pp., Cont.-in-part of U.S. Ser. No. 831,284,

abandoned.

CODEN: USXXAM

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 9

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6268392	B1	20010731	US 1998-37308	19980309

				_						
	CA	2506703		AA	19970918	CA	1997-2506703		19970311	
	$\mathbf{EP}$	1440972		A1	20040728	EP	2004-10088		19970311	
		R: AT, B	E, CH,	DE,	DK, ES, FR,	GB, GI	R, IT, LI, LU,	NL, S	E, PT, IE,	FI
	ΑU	761249		B2	20030529	AU	2000-53394		20000816	
	US	6420417		В1	20020716	US	2000-676466		20000929	
	US	2003171426		A1	20030911	US	2002-76091		20020215	
	US	6642268		B2	20031104					
	US	2004157915		A1	20040812	US	2003-620460		20030717	
	US	6943189		B2	20050913					
PRIO	RIT	APPLN. IN	FO.:			US	1994-305526	A2	19940912	
						US	1995-517051	A1	19950821	
						US	1996-13119P	P	19960311	
						US	1997-40660P	P	19970311	
						US	1997-816065	A2	19970311	
						US	1997-831284	B2	19970331	
						AU	1997-23266	A3	19970311	
						CA	1997-2248586	A3	19970311	
						EP	1997-915976	A3	19970311	
						US	1998-37308	A3	19980309	
						US	2000-676466	A3	20000929	
						US	2002-76091	A1	20020215	

OTHER SOURCE(S):

MARPAT 135:137410

GI

Title compds. [I; R = H or 1-4 of alkyl, alkenyl, alkynyl, acyloxy, aryl, aralkyl, halo, etc.; R1, R2 = H, (substituted) alkyl, alkenyl, alkynyl, haloalkyl, alkylaryl, alkoxy, dialkylamino, etc.; R1R2C = cycloalkylidene; R3, R4 = H, alkyl, alkenyl, alkynyl, acyloxy, aryl, heterocyclyl, heteroaryl, etc.; R3R4 = O, S, NOR11, etc.; R11 = H, alkyl, alkenyl, alkynyl, aryl, etc.; R5, R6 = H, alkyl, alkenyl, alkynyl, aryl, cycloalkyl, heterocyclyl, etc.; R7, R8 = H, alkyl; Z = SOO-2], were prepared A composition comprising an ileal bile acid transport inhibitor and an HMG Co-A reductase inhibitor is claimed. Thus, title compound (II) (preparation via 2-mercapto-4-methoxybenzophenone given) at 0.2% as an ileal perfusion in guinea pigs reduced HDL cholesterol from 89 mg% to 76 mg%.

IC ICM A61K031-38

ICS C07D337-12

INCL 514431000

CC 27-21 (Heterocyclic Compounds (One Hetero Atom))

Section cross-reference(s): 1

IT 178678-22-3P 178678-23-4P 178678-24-5P 178678-25-6P 178678-26-7P 178678-27-8P 178678-28-9P 178678-29-0P 178678-30-3P 178678-31-4P 178678-34-7P 178678-35-8P 178678-36-9P 178678-37-0P 178678-33-6P 178678-38-1P 178678-39-2P 178678-40-5P 178678-43-8P 178678-44-9P 178678-48-3P 178678-51-8P 178678-52-9P ⇒178678-53-0P 178678-45-0P 178897-95-5P 178897-96-6P 178897-97-7P 178897-98-8P 178678-54-1P 178897-99-9P 178898-00-5P 178898-01-6P 178898-02-7P 178898-03-8P

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178898-04-9P
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                                                              197372-69-3P
197372-70-6P
               197372-71-7P
                               197372-72-8P
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197372-75-1P
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                                                              197373-70-9P
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                                              197373-77-6P
                                                              197373-78-7P
197373-79-8P
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                               197373-83-4P
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                                                              197373-87-8P
197373-93-6P
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197374-01-9P
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                               197374-03-1P 197374-04-2P
                               197374-09-7P
197374-06-4P
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                                              197374-10-0P
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                               197374-16-6P
197374~13-3P
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                                              197374-17-7P
                                                              197374-18-8P
197374-19-9P
               197374-20-2P
                               197374-21-3P
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197374-30-4P
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197374-60-0P
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                               197374-63-3P
                                              197374-65-5P
                                                              197374-66-6P
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                                              197374-72-4P
                                                              197374-73-5P
197374-74-6P
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                               197374-76-8P
                                              197374-77-9P
                                                              197374-78-0P
197374-79-1P
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                                              197374-82-6P
                                                              197374-83-7P
197374-85-9P
               197374-86-0P
                               197374-87-1P
                                              197374-88-2P
                                                              197374-89-3P
197374-90-6P
               197374-93-9P
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                                              197374-95-1P
                                                              197374~96-2P
197374-97-3P
               197374-98-4P
                               197374-99-5P
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                                                              197375-01-2P
197375-02-3P
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197375-07-8P
               197375-08-9P
                                              197375-10-3P
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study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); USES (Uses)
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        combination therapy with HMG Co-A reductase inhibitors)
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        (preparation of ileal bile acid transport inhibiting benzothiepines for
        combination therapy with HMG Co-A reductase inhibitors)
IT
     197373-37-8P 197374-04-2P 197374-59-7P
     197375-96-5P 197376-55-9P 197384-36-4P
     213312-84-6P 213313-20-3P
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
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(preparation of ileal bile acid transport inhibiting benzothiepines for

BIOL (Biological study); PREP (Preparation); USES (Uses)

combination therapy with HMG Co-A reductase inhibitors)

RN 197373-37-8 HCAPLUS

CN 1-Propanesulfonic acid, 3-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197374-04-2 HCAPLUS

CN 2-Propenamide, N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197374-59-7 HCAPLUS

CN Propanamide, 3-bromo-N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197384-36-4 HCAPLUS

CN 1-Hexanaminium, 6-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-6-oxo-, rel-, salt with trifluoroacetic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 197384-35-3 CMF C38 H62 N3 O4 S

Relative stereochemistry.

CM 2

CRN 14477-72-6 CMF C2 F3 O2

RN 213312-84-6 HCAPLUS

CN 1-Pentanaminium, 5-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-5-oxo-, iodide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

• I.

RN 213313-20-3 HCAPLUS

CN Benzenaminium, 3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]-N,N,N-triethyl-, bromide, rel-(9CI) (CA INDEX NAME)

● Br-

IT 197373-50-5P 197373-51-6P 213312-74-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of ileal bile acid transport inhibiting benzothiepines for combination therapy with HMG Co-A reductase inhibitors)

RN 197373-50-5 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-5-(3-nitrophenyl)-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197373-51-6 HCAPLUS

CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

RN 213312-74-4 HCAPLUS

CN Pentanamide, 5-chloro-N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

REFERENCE COUNT: 76 THERE ARE 76 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 10 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:590035 HCAPLUS

DOCUMENT NUMBER: 133:193089

TITLE: Preparation of substituted 5-aryl-benzothiepines as

ileal bile acid transport and taurocholate uptake

inhibitors

INVENTOR(S): Lee, Len F.; Banerjee, Shyamal C.; Huang, Horng-chih;

Li, Jinglin J.; Miller, Raymond E.; Reitz, David B.;

Tremont, Samuel J.

PATENT ASSIGNEE(S): G.D. Searle and Co., USA

SOURCE: U.S., 191 pp., Cont.-in-part of U.S. Ser. No.

109,551.

CODEN: USXXAM

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 9
PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
HC 6107404	7 20000022	US 1999-275463 CA 1997-2506703	10000334
CD 2506703	A 20000822	CN 1997-2506703	19990324
FD 1440972	A1 20040728	EP 2004-10088	19970311
		GB, GR, IT, LI, LU, NL,	
US 5994391	A 19991130	US 1998-109551	19980702
EP 1331225	A1 20030730	US 1998-109551 EP 2003-5459	19981216
		GB, GR, IT, LI, LU, NL,	
CA 2336315	AA 20000113	CA 1999-2336315	19990629
WO 2000001687	A1 20000113	CA 1999-2336315 WO 1999-US12828	19990629
W: AL, AM, AT	T, AU, AZ, BA, BB,	BG, BR, BY, CA, CH, CN,	CU, CZ, DE,
		GH, GM, HR, HU, ID, IL,	
KG, KP, KF	R, KZ, LC, LK, LR,	LS, LT, LU, LV, MD, MG,	MK, MN, MW,
MX, NO, NZ	Z, PL, PT, RO, RU,	SD, SE, SG, SI, SK, SL,	TJ, TM, TR,
	G, US, UZ, VN, YU,		
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	A, GN, GW, ML, MR,		
AU 9948202,	A1 20000124 B2 20031030	AU 1999-48202	19990629
AU 766957	B2 20031030		
		EP 1999-931769	19990629
EP 1091953			
		GB, GR, IT, LI, LU, NL,	SE, MC, PT,
IE, SI, LT	T, LV, FI, RO		
TR 200100824	T2 20010723	TR 2001-200100824	19990629
BR 9911737	A 20011211	BR 1999-11737	19990629
EE 200100002	A . 20020617	EE 2001-2	19990629
JP 2002519418	T2 20020702	JP 2000-558091	19990629
NZ 509621	A 20030829	NZ 1999-509621	19990629
AT 256122	E 20031215	AT 1999-931769	19990629
PT 1091953	T 20040430	PT 1999-931769	19990629
ES 22133/3	T3 20040816	ES 1999-931769	19990629
EP 1466911	A2 20041013 A3 20050907	TR 2001-200100824 BR 1999-11737 EE 2001-2 JP 2000-558091 NZ 1999-509621 AT 1999-931769 PT 1999-931769 ES 1999-931769 EP 2003-26649	19990629
DI I-00011	A3 20030707		
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110 6262277	R1 20030128	SG 2002-200207701 US 1999-443403	19990029
TW 229670	B1 20010717	TW 1000-00111202	20000107
AII 761249	. B2 20030521	TW 1999-88111293 AU 2000-53394	20000107
NO 200100016	A 20030329	NO 2001-16	20000010
ZA 2001000018	A 20010302	ZA 2001-28	20010102
HR 2001000026	A1 20010723	HR 2001-4	20010102
BG 105206	A 20011231	BG 2001-105206	20010102
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AU 2004200346	A1 20040226	AU 2004-200346	20040130
JP 2004203891	A2 20040722	JP 2004-50473	20040225
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JP 2004359694	A2 20041224	JP 2004-227034	20040803

PRIORITY	APPLN	INFO. :	

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US	2002-68297	A3	20020208

III

OTHER SOURCE(S):

MARPAT 133:193089

$$(R?)_{q} \xrightarrow{(O)_{n}} R^{7}$$

$$R^{8}$$

$$R^{1}$$

$$R^{2}$$

$$R^{6}$$

$$R^{5}$$

$$R^{4}$$

$$R^{3}$$

$$R^{3}$$

The title compds. (I) [wherein q = 1-4; n = 2; R1 and R2 = independently H or (un)substituted (halo)alkyl, alkenyl, alkynyl, alkylaryl, arylalkyl, alkoxy(alkyl), dialkylamino, alkylthio, (polyalkyl)aryl, or cycloalkyl; or R1 and R2 taken together with the atoms to which they are attached = cycloalkyl; R3 and R4 = independently H, alkyl, alkenyl, alkynyl, acyloxy, aryl, heterocyclyl, OR9, NR9R10, SR9, S(O)R9, SO2R9, or SO3R9; R9 and R10 = independently H, (cyclo)alkyl, alkenyl, alkynyl, aryl(alkyl), acyl, heterocyclyl, or ammoniumalkyl; or R3 and R4 together = :O, :NOR11, :S, :NNR11R12, :NR9, or :CR11R12; R11 and R12 = independently H, (cyclo)alkyl, alkenyl, alkynyl, aryl(alkyl), heterocyclyl, carboxylalkyl, carboalkoxyalkyl, cyanoalkyl, OR9, NR9R10, SR9, S(O)R9, SO2R9, SO3R9, CO2R9, CN, halo, oxo, or CONR9R10; R5 = substituted aryl; R6 = H; R7 and R8 = independently H or alkyl; Rx = independently H or (un)substituted

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(cyclo)alkyl, alkenyl, alkynyl, polyalkyl, acyloxy; aryl(alkyl),
    halo(alkyl), (quaternary) heterocyclyl, (quaternary) heteroaryl,
    polyether, alkoxy, amino, alkylthio, NO2, carboxy, carbamido, etc.] where
    prepared for the prophylaxis and treatment of hyperlipidemic conditions,
     such as those associated with atherosclerosis or hypercholesterolemia. Thus,
    KOBu-t was added to a solution of 2-((2-benzyl-5-
    methoxyphenylsulfonyl)methyl)-2-ethylhexanal (preparation given) and dry THF
     cooled to -1.6°C to give, after workup, II and III (96% combined
     yield). The isomers were separated upon recrystn. II inhibited IBAT-mediated
     uptake of [14C]-taurocholate in H14 cells with an IC50 of 0.1 \mu M and
     reduced serum cholesterol from 143 mg (7%) to 126 mg (2%) compared to
     control in cholesterol-fed hamsters in a 14-day test. In vitro
     taurocholate uptake assay data are included for nearly 600 compds. of the
     invention.
     C07D337-00; C07D487-00; A61K031-38; A61K031-495
INCL 549009000
     27-21 (Heterocyclic Compounds (One Hetero Atom))
     Section cross-reference(s): 1
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     (Reactant or reagent); USES (Uses)
        (hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by
        cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as
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                                   178897-95-5P
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                                                                  178897-99-9P
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197374-00-8P

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197374-02-0P

197374-03-1P

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197375-26-1P
               197375-28-3P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); USES (Uses)
   (hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by
   cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as
   ileal bile acid transport and taurocholate uptake inhibitors)
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197377-75-6P

197377-76-7P

197377-77-8P

197377-78-9P

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                   289038-29-5P
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     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); USES (Uses)
        (hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by
       cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as
        ileal bile acid transport and taurocholate uptake inhibitors)
                   289038-34-2P 289038-35-3P 289038-36-4P
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     289038-33-1P
     289038-37-5P 289038-38-6P
                               289038-39-7P
                                              289038-40-0P
                   289038-42-2P 289038-43-3P
     289038-41-1P
                                               289038-44-4P
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                   289056-45-7P
                                289056-46-8P
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); USES (Uses)
        (hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by
       cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as
       ileal bile acid transport and taurocholate uptake inhibitors)
IT
     197373-50-5P 197373-51-6P 289037-96-3P
     289037-98-5P
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT
     (Reactant or reagent); USES (Uses)
        (hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by
       cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as
       ileal bile acid transport and taurocholate uptake inhibitors)
RN
     197373-50-5 HCAPLUS
     1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-5-(3-
CN
     nitrophenyl)-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)
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RN 197373-51-6 HCAPLUS
CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3,3-dibutyl-7-(dimethylamino)2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 289037-96-3 HCAPLUS
CN Carbamic acid, [3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-,
3-chloropropyl ester, rel- (9CI) (CA INDEX NAME)

RN 289037-98-5 HCAPLUS

Urea, N-[3-(chloromethyl)phenyl]-N'-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

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IT
     197373-37-8P 197373-54-9P 197374-04-2P
     197374-59-7P 197375-96-5P 197376-42-4P
     197376-55-9P 197384-36-4P 289037-97-4P
     289037-99-6P 289038-26-2P 289038-27-3P
     289038-28-4P 289038-35-3P 289038-36-4P
     289038-37-5P 289038-38-6P 289038-43-3P
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); USES (Uses)
        (hypolipemic agent; preparation of substituted 5-aryl-benzothiepines by
        cyclization of 2-((2-benzyl- and 2-benzoylphenylthio)methyl)alkanals as
        ileal bile acid transport and taurocholate uptake inhibitors)
RN
     197373-37-8 HCAPLUS
CN
    1-Propanesulfonic acid, 3-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-
```

2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197373-54-9 HCAPLUS

CN 1-Pentanaminium, 5-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-5-oxo-, rel-, salt with trifluoroacetic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 197373-53-8 CMF C37 H60 N3 O4 S

Relative stereochemistry.

CM 2

CRN 14477-72-6 CMF C2 F3 O2

197374-04-2 HCAPLUS RN

2-Propenamide, N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-CN tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN

197374-59-7 HCAPLUS Propanamide, 3-bromo-N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-CNtetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

RN 197375-96-5 HCAPLUS

, <u>`</u>

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-5-[2-(dimethylamino)phenyl]-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel-(9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197376-42-4 HCAPLUS

CN Benzenaminium, 3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]-N,N,N-trimethyl-, iodide, rel-(9CI) (CA INDEX NAME)

Relative stereochemistry.

• I-

RN 197376-55-9 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-5-[3-(dimethylamino)phenyl]-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel-(9CI) (CA INDEX NAME)

RN 197384-36-4 HCAPLUS

CN 1-Hexanaminium, 6-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-6-oxo-, rel-, salt with trifluoroacetic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 197384-35-3 CMF C38 H62 N3 O4 S

Relative stereochemistry.

CM 2

CRN 14477-72-6 CMF C2 F3 O2

RN 289037-97-4 HCAPLUS

CN 4-Aza-1-azoniabicyclo[2.2.2]octane, 1-[3-[[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]carbonyl]oxy]propyl]-, chloride, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

$$n-Bu$$
 $n-Bu$ 
 $R$ 
 $R$ 
 $NMe_2$ 
 $HN$ 
 $O$ 
 $(CH_2)_3$ 

● Cl -

RN 289037-99-6 HCAPLUS

CN 4-Aza-1-azoniabicyclo[2.2.2]octane, 1-[[3-[[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]carbonyl]amino]phenyl]methyl]-, chloride, rel- (9CI) (CA INDEX NAME)

● Cl -

RN 289038-26-2 HCAPLUS

CN Pyridinium, 2-[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]methyl]-1-ethyl-, iodide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

DI-

RN 289038-27-3 HCAPLUS

CN Pyridinium, 3-[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]methyl]-1-ethyl-, iodide, rel- (9CI) (CA INDEX NAME)

• I-

Relative stereochemistry.

• I -

RN 289038-35-3 HCAPLUS
CN Pyridinium, 2-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-1-ethyl-, iodide, rel- (9CI) (CA INDEX NAME)

● T~

RN 289038-36-4 HCAPLUS
CN Pyridinium, 3-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-1-ethyl-, iodide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

• I.

RN 289038-37-5 HCAPLUS
CN Benzenemethanaminium, 4-[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]sulfonyl]-N,N,N-triethyl-, iodide, rel- (9CI) (CA INDEX

NAME)

Relative stereochemistry.

• I-

RN 289038-38-6 HCAPLUS Pyridinium, 4-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-CNtetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-1-ethyl-, iodide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN289038-43-3 HCAPLUS CN Benzenaminium, 3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]-N,N,N-triphenyl-, bromide, rel-(9CI) (CA INDEX NAME)

Relative stereochemistry.

● Br-

REFERENCE COUNT:

THERE ARE 56 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 11 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

56

ACCESSION NUMBER:

2000:456922 HCAPLUS

DOCUMENT NUMBER:

133:94515

TITLE:

Combinations for cardiovascular indications

INVENTOR(S):

Keller, Bradley T.; Reitz, David B.; Schuh, Joseph R.;

Sikorski, James A.; Tremont, Samuel J.; Lappe, Rodney

W.

PATENT ASSIGNEE(S):

G.D. Searle and Co., USA

SOURCE:

PCT Int. Appl., 248 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE		
WO 2000038725	A1 20000706	WO 1999-US27946	19991217		
W: AE, AL, AM,	AT, AU, AZ, BA,	BB, BG, BR, BY, CA, CH,	CN, CR, CU,		
CZ, DE, DK,	DM, EE, ES, FI,	GB, GD, GE, GH, GM, HR,	HU, ID, IL,		
IN, IS, JP,	KE, KG, KP, KR,	KZ, LC, LK, LR, LS, LT,	LU, LV, MA,		
MD, MG, MK,	MN, MW, MX, NO,	NZ, PL, PT, RO, RU, SD,	SE, SG, SI,		
SK, SL, TJ,	TM, TR, TT, TZ,	UA, UG, US, UZ, VN, YU,	ZA, ZW, AM,		
AZ, BY, KG,	KZ, MD, RU, TJ,	TM			
RW: GH, GM, KE,	LS, MW, SD, SL,	SZ, TZ, UG, ZW, AT, BE,	CH, CY, DE,		
DK, ES, FI,	FR, GB, GR, IE,	IT, LU, MC, NL, PT, SE,	BF, BJ, CF,		
CG, CI, CM,	GA, GN, GW, ML,	MR, NE, SN, TD, TG			
CA 2356515	AA 20000706	CA 1999-2356515	19991217		
EP 1140187	A1 20011010	EP 1999-965901	19991217		
EP 1140187	B1 20030903				

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                                            20020129 BR 1999-16564
                                 T2 20021008 JP 2000-590676
A1 20030319 EP 2002-25631
      JP 2002533411
      OF 2002533411
EP 1293211
           R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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A1 20030820 EP 2003-9706
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            R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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      EP 1340508
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      EP 1340510
            R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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                                   A1 20030910 EP 2003-11146
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                                  E 20030915 AT 1999-965901 19991217
A1 20031022 EP 2003-16600 19991217
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EP 1354604
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            R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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      NZ 512532 A
                                           20031219 NZ 1999-512532
                                         20040130 PT 1999-965901
      PT 1140187
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      ES 2207330
AU 779264
ZA 2001005056
     ES 2207330 T3 20040516 ES 1999-965901
AU 779264 B2 20050113 AU 2000-21577
ZA 2001005056 A 20020620 ZA 2001-5056
ZA 2001005059 A 20020620 ZA 2001-5059
ZA 2001005061 A 20020620 ZA 2001-5061
ZA 2001005062 A 20020828 ZA 2001-5062
ZA 2001005060 A 20020920 ZA 2001-5060
NO 2001003157 A 20010822 NO 2001-3157
US 2003166720 A1 20030904 US 2002-200600
US 2003203892 A1 20031030 US 2002-200599
US 2003109558 B2 20050510
US 6890958
B2 20050510
                                                                                            19991217
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      US 6890958 B2 20050510
US 2003125316 A1 20030703 US 2002-245507
US 2004058908 A1 20040325 US 2002-266743
US 2004029845 A1 20040212 US 2003-373180
US 2004028644 A1 20040212 US 2003-412694
US 2004048846 A1 20040311 US 2003-652306
                                                                                             20020918
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                                                            US 2003-412694 20030414

US 2003-652306 20030902

US 1998-113955P P 19981223

US 1999-142603P P 19990707
PRIORITY APPLN. INFO.:
                                                             US 1999-142616P
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                                                             US 1999-142682P
                                                                                        P 19990707
                                                             US 1999-142684P
                                                                                        P 19990707
                                                             US 1999-143043P
                                                                                        P 19990707
                                                             US 1999-143047P
                                                                                        P 19990707
                                                             US 1999-143047P P 19990707
US 1999-143550P P 19990713
EP 1999-965035 A3 19991217
EP 1999-965899 A3 19991217
EP 1999-965900 A3 19991217
EP 1999-965902 A3 19991217
EP 1999-965903 A3 19991217
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A3 19991217
EP 1999-967140
                 A3 19991217
A3 19991217
US 1999-465642
US 1999-466413
US 1999-466415
                 A3 19991217
                 B1 19991217
US 1999-466466
US 1999-466469
                 A3 19991217
US 1999-466470
                  A3 19991217
US 1999-466592
                  A3 19991217
US 1999-466596
                  B3 19991217
WO 1999-US27946
                 W 19991217
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- The present invention provides combinations of cardiovascular therapeutic compds. for the prophylaxis or treatment of cardiovascular disease including hypercholesterolemia and atherosclerosis. Combinations disclosed include an ileal bile acid transport inhibitor combined with a cholesteryl ester transport protein (CETP) inhibitor, a fibric acid derivative, a nicotinic acid derivative, a microsomal triglyceride transfer protein inhibitor, a cholesterol absorption antagonist, a phytosterol, a stanol, an antihypertensive agent, or others. Further combinations include a CETP inhibitor with a fibric acid derivative, a nicotinic acid derivative, a bile acid sequestrant, a microsomal triglyceride transfer protein inhibitor, a cholesterol absorption antagonist, or others.
- IC ICM A61K045-06
  - ICS A61K031-55; A61K031-585; A61P009-00; A61K031-575
- CC 63-6 (Pharmaceuticals)
  - Section cross-reference(s): 1

280106-13-0D, enantiomers

IT 52-01-7, Spironolactone 59-67-6D, Nicotinic acid, derivs Cholestanol 83-45-4, Fucostanol 360-68-9, Coprostanol 474-60-2, 516-95-0, Epicholestanol 943-45-3D, Fibric acid, derivs Campestanol 23288-49-5, Probucol 55529-51-6, Clionastanol 96829-58-2, 4651-51-8 Orlistat 107724-20-9, Eplerenone 114798-26-4, Losartan 138126-65-5, Stigmastanol 163222-33-1 178961-24-5D, enantiomers 197372-90-0D, enantiomers 197373-37-8D, enantiomers 197373-42-5D, enantiomers 197373-57-2D, enantiomers 229307-33-9D, enantiomers 280105-79-5D, enantiomers 280105-80-8D, enantiomers 280105-82-0D, enantiomers 280105-83-1D, enantiomers 280105-84-2D, enantiomers 280105-85-3D, enantiomers **280105-86-4D**, enantiomers 280105-88-6D, enantiomers 280105-89-7D, enantiomers **280105-90-0D**, enantiomers 280105-91-1D, enantiomers 280105-92-2D, enantiomers 280105-94-4D, enantiomers 280105-95-5D, enantiomers 280105-96-6D, enantiomers 280105-97-7D, enantiomers 280105-98-8D, enantiomers 280105-99-9D, enantiomers 280106-00-5D, enantiomers 280106-01-6D, enantiomers 280106-02-7D, enantiomers 280106-03-8D, enantiomers 280106-04-9D, enantiomers 280106-05-0D, enantiomers 280106-06-1D, enantiomers 280106-08-3D, enantiomers 280106-09-4D, enantiomers 280106-10-7D, enantiomers 280106-11-8D, enantiomers 280106-12-9D, enantiomers

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(combinations for cardiovascular agents for treatment of cardiovascular indications)

IT 197373-37-8D, enantiomers 280105-86-4D, enantiomers
280105-88-6D, enantiomers 280105-89-7D, enantiomers
280105-90-0D, enantiomers 280105-97-7D, enantiomers
280105-98-8D, enantiomers

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(combinations for cardiovascular agents for treatment of cardiovascular indications)

RN 197373-37-8 HCAPLUS

CN 1-Propanesulfonic acid, 3-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 280105-86-4 HCAPLUS

CN 1-Pentanaminium, 5-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-5-oxo-, chloride, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

● c1-

RN 280105-88-6 HCAPLUS

CN Pyridinium, 4-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-1-

methyl-, rel-, salt with 4-methylbenzenesulfonic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 280105-87-5 CMF C32 H44 N3 O3 S

Relative stereochemistry.

CM 2

CRN 16722-51-3 CMF C7 H7 O3 S

RN 280105-89-7 HCAPLUS

CN 4-Aza-1-azoniabicyclo[2.2.2]octane, 1-[[4-[[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]methyl]phenyl]methyl]-, chloride, rel- (9CI) (CA INDEX NAME)

. 3 4.

Ocl-

$$\begin{array}{c} \text{NMe}_2 \\ \text{NMe}_2 \\ \text{MeO} \\ \hline \end{array} \\ \begin{array}{c} \text{CH}_2 - \text{CH}_2 - \text{OH}_2 - \text{NH} - \text{C-NH} \\ \text{O} \\ \end{array}$$

RN 280105-97-7 HCAPLUS

CN 1-Pentanaminium, 5-[[3-[(3R,4S,5S)-3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-5-oxo-, chloride, rel- (9CI) (CA INDEX NAME)

● Cl~

RN 280105-98-8 HCAPLUS

CN 1-Propanesulfonic acid, 3-[[3-[(3R,4S,5S)-3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 12 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

2

ACCESSION NUMBER:

1999:795803 HCAPLUS

DOCUMENT NUMBER:

132:35625

TITLE:

Amino acid containing benzo[b]thiepine 1,1-dioxide

derivatives as hypolipemic agents

INVENTOR(S):

Frick, Wendelin; Enhsen, Alfons; Glombik, Heiner;

Heuer, Hubert

PATENT ASSIGNEE(S):

Hoechst Marion Roussel Deutschland G.m.b.H., Germany

SOURCE: PCT Int. Appl., 28 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent German

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.			KIND	DATE	APPLICATION NO.		
	0 9964410			A1		WO 1999-EP3701	19990528
WO		ΔT.	ΔM			BB, BG, BR, BY, CA, CH,	
	W. AE,	DK	EE.	ES FI	GB GD	GE, GH, GM, HR, HU, ID,	II. IN. IS.
	.TD	KE,	KG.	KP KR	, GB, GB,	LK, LR, LS, LT, LU, LV,	MD. MG. MK.
						RO, RU, SD, SE, SG, SI,	
						VN, YU, ZA, ZW	411, 42, 11,
						SZ, UG, ZW, AT, BE, CH,	CY, DE, DK,
	ES.	FI.	FR.	GB. GR	. IE. IT.	LU, MC, NL, PT, SE, BF,	BJ, CF, CG,
						NE, SN, TD, TG	
DE	19825804		,	A1		DE 1998-19825804	19980610
	19825804			C2	20000824		
CA	2334775			AA	19991216	CA 1999-2334775	19990528
AU	9945019			A1	19991230	AU 1999-45019	19990528
					20021010		
EP	1086092			A1	20010328	EP 1999-927784	19990528
EP	1086092			B1	20021113		
	R: AT,	BE,	CH,	DE, DK	, ES, FR,	GB, GR, IT, LI, LU, NL,	
BR	9912188			Α	20010410	BR 1999-12188 TR 2000-200003634	19990528
TR	20000363	4		T2	20010621	TR 2000-200003634	
JP	20025174				20020618		
AΤ	227715			E	20021115	AT 1999-927784	
ES	2182535			<b>T</b> 3	20030301	ES 1999-927784	19990528
	1086092			T	20030331	PT 1999-927784	
	2215001			C2	20031027		19990528
	1127497			В	20031112	CN 1999-807171	19990528
	20000363	2		T2	20010420		
	1086113			T	20040630		19990529
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	1036799	T.1700		AI	20040402		20011106
CTORT'T	Y APPLN.	TNFO	.:			DE 1998-19825804 AU 1997-23266	A 19980610 A3 19970311
							W 19990528
							A1 19990920
uren e	OTTOCE (C) .			מממגוש	132:3562		A1 13330320
	OURCE(S):			PIARPA	132:3362	<b>.</b>	
[							

<sup>\*</sup> STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

Title compds. such as I (mixture of diastereoisomers) were prepared as AB hypolipemic agents. Thus, I was prepared in 2 sequences from racemic II and Fmoc-D-lys(Boc)-OH, followed by removal of the Fmoc group with Et2NH. I was ≥20 times more active than 3 analogous comparison substances in

tests of fecal separation of 14C-taurocholic acid in rats.

IC ICM C07D337-08

ICS C07K005-068; A61K031-38

CC 27-21 (Heterocyclic Compounds (One Hetero Atom))
 Section cross-reference(s): 1

IT 252372-02-4P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(amino acid containing benzo[b]thiepine 1,1-dioxide derivs. as hypolipemic agents)

IT 92122-45-7 252047-42-0

RL: RCT (Reactant); RACT (Reactant or reagent)
 (amino acid containing benzo[b]thiepine 1,1-dioxide derivs. as hypolipemic
agents)

IT 252372-00-2P 252372-01-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(amino acid containing benzo[b]thiepine 1,1-dioxide derivs. as hypolipemic agents)

IT 252372-02-4P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(amino acid containing benzo[b]thiepine 1,1-dioxide derivs. as hypolipemic agents)

RN 252372-02-4 HCAPLUS

CN D-Lysinamide, N6-[(1,1-dimethylethoxy)carbonyl]-D-lysyl-N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-N6-[(1,1-dimethylethoxy)carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 252047-42-0

RL: RCT (Reactant); RACT (Reactant or reagent)
 (amino acid containing benzo[b]thiepine 1,1-dioxide derivs. as hypolipemic
 agents)

RN 252047-42-0 HCAPLUS

CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-, 1,1-dioxide, (3R,4S,5S)-rel- (9CI) (CA INDEX NAME)

IT 252372-00-2P 252372-01-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(amino acid containing benzo[b]thiepine 1,1-dioxide derivs. as hypolipemic agents)

RN 252372-00-2 HCAPLUS

CN Carbamic acid, [(1R)-1-[[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]carbonyl]-5-[[(1,1-dimethylethoxy)carbonyl]amino]pentyl]-, 9H-fluoren-9-ylmethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 252372-01-3 HCAPLUS

CN Carbamic acid, [(5R)-5-amino-6-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-6-oxohexyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

$$NMe_2$$
 $NH_2$ 
 $NH_$ 

REFERENCE COUNT: THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS 1 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 13 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1999:795802 HCAPLUS

DOCUMENT NUMBER:

132:22884

TITLE:

Preparation of benzothiepine-1,1-dioxides as

hypolipemics

INVENTOR(S):

Frick, Wendelin; Enhsen, Alfons; Glombik, Heiner;

Heuer, Hubert

PATENT ASSIGNEE(S):

Hoechst Marion Roussel Deutschland G.m.b.H., Germany

SOURCE:

PCT Int. Appl., 30 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.					DATE		APPLICATION NO.							DATE				
				-														
WO 9964409			A2		1999	1216	WO 1999-EP3743							19990529				
WO	9964	409			A3		2000	0302										
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		JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	
		MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	
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		CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG						
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TR	2000	0363	4		T2	2 20010621			TR 2000-200003634							19990528		
ES	2182	535			Т3		2003	0301	ES 1999-927784							19990528		
PT	1086	092			$\mathbf{T}$		2003	0331	PT 1999-927784							19990528		
CN	1127	497			В		2003	1112	CN 1999-807171							19990528		
CA	2334	773			AA		1999	1216										
ΑU	9945	031			A1		1999	1230		AU 1	1999-	4503	1		1	9990	529	
ΑU	7526	33			B2		2002	0926										
EP	1086	113			A2		2001	0328		EP 1	1999-	9278	02		1	9990	529	

EP 1086113	B1	20040211		
R: AT, BE,	CH, DE,	DK, ES, FR,	GB, GR, IT, LI, LU, NL,	SE, PT, IE, FI
TR 200003632	T2	20010420	TR 2000-200003632	
JP 2002517490	T2	20020618	JP 2000-553418	19990529
JP 3374129	B2	20030204		
NZ 508681	A	20020628	NZ 1999-508681	19990529
RU 2220141	C2	20031227	RU 2001-101499	19990529
AT 259372	E	20040215	AT 1999-927802	19990529
PT 1086113	T	20040630	PT 1999-927802	19990529
IL 140078	A1	20040831	IL 1999-140078	19990529
ES 2215387	Т3	20041001	ES 1999-927802	19990529
BR 9911123	Α	20060103	BR 1999-11123	19990529
US 6221897	B1	20010424	US 1999-398315	19990920
AU 761249	B2	20030529	AU 2000-53394	20000816
ZA 2000007060	A	20010718	ZA 2000-7060	20001130
ZA 2000007061	A	20010718	ZA 2000-7061	20001130
NO 2000006251	Α	20010207	NO 2000-6251	20001208
US 2002045583	A1	20020418	US 2001-773772	20010202
US 6441022	B2	20020827		
HK 1039490	A1	20041210	HK 2001-107746	20011106
US 2003017996	A1	20030123	US 2002-201050	20020724
US 6642269	B2	20031104		
US 2004087648	A1	20040506	US 2003-606771	20030627
US 7019023	B2	20060328		
PRIORITY APPLN. INFO.	. <b>:</b>		DE 1998-19825804	A 19980610
				P 19960311
				A3 19970311
				W 19990529
				Al 19990920
				A1 20010202
			US 2002-201050	A1 20020724

OTHER SOURCE(S):

MARPAT 132:22884

GI

$$\mathbb{R}^{4\mathbb{R}^{5}N}$$

$$\mathbb{R}^{0}$$

$$\mathbb{R}^{1}$$

$$\mathbb{R}^{2}$$

AB Title compds. [I; R = C6H4NHZR3; R1,R4,R5 = Me, Et, Pr, Bu; R2 = H, OH, amino(alkyl); R3 = sugar residue; Z = bond, carbonyl(alkylene), CONH, etc.] were prepared Thus, I [R = C6H4(NHR')-3, R1 = Et, R2 = OH, R4 = R5 = Me](II; R' = H) was amidated by penta-O-acetyl-D-gluconic acid and the product deprotected to give II (R' = gluconoyl) as a mixture of diastereomers. Data for biol. activity of I were given.

IC ICM C07D337-00

CC 27-21 (Heterocyclic Compounds (One Hetero Atom))
 Section cross-reference(s): 1

252208-69-8P 252208-70-1P 252208-71-2P

IT 252047-36-2P 252047-37-3P 252047-38-4P 252047-39-5P 252047-40-8P 252047-41-9P 252208-66-5P 252208-67-6P 252208-68-7P

Searched by Paul Schulwitz 571-272-2527

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of benzothiepine-1,1-dioxides as hypolipemics) IT 488-43-7, D-Glucamine 2432-99-7, 11-Aminoundecanoic acid 17430-71-6, Penta-O-acetyl-D-gluconic acid 53555-69-4 252047-42-0 252047-43-1 RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of benzothiepine-1,1-dioxides as hypolipemics) IT 252047-36-2P 252047-37-3P 252047-38-4P 252047-39-5P 252047-40-8P 252047-41-9P 252208-66-5P 252208-67-6P 252208-68-7P 252208-69-8P 252208-70-1P 252208-71-2P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of benzothiepine-1,1-dioxides as hypolipemics) RN252047-36-2 HCAPLUS CN D-Glucitol, 1-[[5-[[3-[(3R,4S,5S)-3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-5-oxopentyl]amino]-1-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 252047-38-4 HCAPLUS

CN D-Gluconamide, N-[3-[3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, 2,3,4,5,6-pentaacetate (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 252047-39-5 HCAPLUS

CN D-Gluconamide, N-[3-[3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]- (9CI) (CA INDEX NAME)

RN 252047-40-8 HCAPLUS

CN D-Glucitol, 1-[[5-[[3-[(3S,4R,5R)-3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-5-oxopentyl]amino]-1-deoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 252047-41-9 HCAPLUS

CN D-Glucitol, 1-deoxy-1-[[5-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-5-oxopentyl]amino]- (9CI) (CA INDEX NAME)

$$n-Bu$$
 $n-Bu$ 
 $n-Bu$ 

RN 252208-66-5 HCAPLUS

CN D-Gluconamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 252208-67-6 HCAPLUS

CN D-Gluconamide, N-[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, 2,3,4,5,6-pentaacetate (9CI) (CA INDEX NAME)

RN 252208-68-7 HCAPLUS

CN D-Gluconamide, N-[11-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-11-oxoundecyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Et 
$$n-Bu$$

NMe2

HN (CH2) 10 OH OH OH OH OH OH OH OH

RN 252208-69-8 HCAPLUS

CN D-Gluconamide, N-[5-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-5-oxopentyl]-, 2,3,4,5,6-pentaacetate (9CI) (CA INDEX NAME)

RN 252208-70-1 HCAPLUS

CN D-Gluconamide, N-[5-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-5-oxopentyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 252208-71-2 HCAPLUS

CN D-Glucitol, 1-[acetyl[5-[[3-[3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-5-oxopentyl]amino]-1-deoxy-, 2,3,4,5,6-pentaacetate (9CI) (CA INDEX NAME)

IT 252047-42-0 252047-43-1

RL: RCT (Reactant); RACT (Reactant or reagent)
 (preparation of benzothiepine-1,1-dioxides as hypolipemics)

RN 252047-42-0 HCAPLUS

CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-, 1,1-dioxide, (3R,4S,5S)-rel-(9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 252047-43-1 HCAPLUS

CN Pentanamide, 5-bromo-N-[3-[(3R,4S,5S)-3-butyl-7-(dimethylamino)-3-ethyl-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

L9 ANSWER 14 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1998:621210 HCAPLUS

DOCUMENT NUMBER: 129:260353

TITLE: Preparation of ileal bile acid transport inhibiting

benzothiepines for combination therapy with HMG Co-A

reductase inhibitors.

INVENTOR(S): Reitz, David B.; Lee, Len F.; Li, Jinglin J.; Huang,

Horng-Chih; Tremont, Samuel J.; Miller, Raymond E.; Baneriee, Shyamal C.; Manning, Robert E.; Glenn, Kevin

C.; Keller, Bradley T.

PATENT ASSIGNEE(S): G.D. Searle and Co., USA; et al.

SOURCE: PCT Int. Appl., 477 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 9

PATENT INFORMATION:

PATENT NO.							DATE											
					-													
WO	9840	375			A2		1998	0917	1	WO 1	998-1	JS379	92		19980310			
WO	9840	375			A3		19981203											
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		DK,	EE,	ES,	FI,	GB,	GE,	GH,	GM,	GW,	HU,	ID,	IL,	IS,	JP,	KE,	KG,	
							LR,											
		NO,	NZ,	PL,	PT,	RO	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	
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	RW:						SD,		UG,	ZW,	AT,	BE,	CH,	DE,	DK,	ES,	FI,	
							LU,											
							SN,			•	•	•	,	-	-	-		
CA	2283								CA 1998-2283575						1	9980	310	
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	7300																	
									EP 1998-910075						10000210			
EP																		
		•		CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LΙ,	ъU,	NL,	SE,	MC,	PI,	
		ΙE,																
										NZ 1998-337830								
BR	9808	013			A		2001	0925		BR 1	998-	8013			19980310			
JP 2002500628				Т2		2002	0108		JP 1	998-	5395	94		19980310				

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PRIORITY APPLN. INFO.:
                                             US 1997-40660P
                                             US 1994-305526
                                                                  B2 19940913
                                             US 1995-517051
                                                                  B1 19950821
                                             US 1996-13119P
                                                                  P 19960311
                                             AU 1997-23266
                                                                  A3 19970311
                                             US 1997-816065
                                                                  B2 19970311
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                                                                  B3 19970331
                                             WO 1998-US3792
                                                                  W 19980310
                                             US 2000-676466
                                                                  A3 20000929
```

OTHER SOURCE(S):

MARPAT 129:260353

GI

$$(R^9)$$
 q  $R^7$   $R^8$   $R^1$   $R^2$   $R^2$   $R^3$   $R^6$   $R^5$   $R^4$   $R^3$   $R^6$   $R^5$   $R^4$   $R^9$   $R^9$ 

Title compds. [I; q = 1-4; n = 0-2; R1, R2 = H, (substituted) alkyl, alkenyl, alkynyl, haloalkyl, alkylaryl, alkoxy, dialkylamino, etc.; R1R2C = cycloalkylidene; R3, R4 = H, alkyl, alkenyl, alkynyl, acyloxy, aryl, heterocyclyl, heteroaryl, etc.; R3R4 = O, S, NOR11, etc.; R11 = H, alkyl, alkenyl, alkynyl, aryl, aralkyl, etc.; R5, R6 = H, alkyl, alkenyl, alkynyl, aryl, cycloalkyl, heterocyclyl, etc.; R7, R8 = H, alkyl; R9 = H, alkyl, alkenyl, alkynyl, acyloxy, aryl, aralkyl, halo, etc.], were prepared A composition comprising an ileal bile acid transport inhibitor and an HMG Co-A reductase inhibitor is claimed. Thus, title compound (II) (preparation via 2-mercapto-4-methoxybenzophenone given) at 0.2% as an ileal perfusion in guinea pigs reduced HDL cholesterol from 89 mg% to 76 mg%.

IC ICM C07D337-00

CC 27-21 (Heterocyclic Compounds (One Hetero Atom))

Section cross-reference(s): 1

IT	178678-22-3P	178678-23-4P	178678-24-5P	178678-25-6P	178678-26-7P
	178678-27-8P	178678-28-9P	178678-29-0P	178678-30-3P	178678-31-4P
	178678-33-6P	178678-34-7P	178678-35-8P	178678-36-9P	178678-37-0P
	178678-38-1P	178678-39-2P	178678-40-5P	178678-43-8P	178678-44-9P
	178678-45-0P	178678-48-3P	178678-51-8P	178678-52-9P	178678-53-0P
	178678-54-1P	178897-95-5P	178897-96-6P	178897-97-7P	178897-98-8P
	178897-99-9P	178898-00-5P	178898-01-6P	178898-02-7P	178898-03-8P
	178898-04-9P	178898-05-0P	197372-66-0P	197372-67-1P	197372-69-3P
	197372-70-6P	197372-71-7P	197372-72-8P	197372-73-9P	197372-74-0P
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BIOL (Biological study); PREP (Preparation); USES (Uses)
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RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);

IT

(preparation of ileal bile acid transport inhibiting benzothiepines for combination therapy with HMG Co-A reductase inhibitors)

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     BIOL (Biological study); PREP (Preparation); USES (Uses)
        (preparation of ileal bile acid transport inhibiting benzothiepines for
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     study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); USES (Uses)
        (preparation of ileal bile acid transport inhibiting benzothiepines for
       combination therapy with HMG Co-A reductase inhibitors)
RN
     197373-37-8 HCAPLUS
CN
     1-Propanesulfonic acid, 3-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-
     2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-
```

Relative stereochemistry.

(CA INDEX NAME)

, rel- (9CI)

RN 197374-04-2 HCAPLUS 2-Propenamide, N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-CN

tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

197374-59-7 HCAPLUS RN

Propanamide, 3-bromo-N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-CNtetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

RN 197375-96-5 HCAPLUS
CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-5-[2-(dimethylamino)phenyl]-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel-(9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197384-36-4 HCAPLUS

CN 1-Hexanaminium, 6-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-6-oxo-, rel-, salt with trifluoroacetic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 197384-35-3 CMF C38 H62 N3 O4 S

Relative stereochemistry.

CM 2

CRN 14477-72-6 CMF C2 F3 O2

RN 213312-84-6 HCAPLUS

CN 1-Pentanaminium, 5-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-5-oxo-, iodide, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

• I -

RN 213313-20-3 HCAPLUS

CN Benzenaminium, 3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]-N,N,N-triethyl-, bromide, rel-(9CI) (CA INDEX NAME)

• Br-

IT 197373-50-5P 197373-51-6P 213312-74-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of ileal bile acid transport inhibiting benzothiepines for combination therapy with HMG Co-A reductase inhibitors)

RN 197373-50-5 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-5-(3-nitrophenyl)-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197373-51-6 HCAPLUS

CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

RN 213312-74-4 HCAPLUS

CN Pentanamide, 5-chloro-N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

## Relative stereochemistry.

L9 ANSWER 15 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:623163 HCAPLUS

DOCUMENT NUMBER: 127:307312

TITLE: Novel benzothiepines having activity as inhibitors of

ileal bile acid transport and taurocholate uptake Reitz, David B.; Lee, Len F.; Li, Jinglin J.; Huang,

Horng-Chih; Tremont, Samuel J.; Miller, Raymond E.;

Banerjee, Shyamal C.

PATENT ASSIGNEE(S): G.D. Searle and Co., USA; Reitz, David B.; Lee, Len

F.; Li, Jinglin J.; Huang, Horng-Chih; Tremont, Samuel

J.; Miller, Raymond E.; Banerjee, Shyamal C.

SOURCE: PCT Int. Appl., 406 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 9

INVENTOR(S):

## PATENT INFORMATION:

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	WO 9733882			A1 19970918			WO 1997-US4076											
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Searched by Paul Schulwitz 571-272-2527

AB Novel benzothiepines I [q = 1-4; n = 0-2; R = H, halo, (un)substitutedalk(en/yn)yl, acyloxy, aryl, heterocyclyl, OH or NH2 or SH or derivs., etc.; R1, R2 = H, (un)substituted and/or heteroatom-replaced alk(en/yn)yl, cycloalkyl, aryl, alkoxy, alkylthio, dialkylamino; or CR1R2 = C3-10 cycloalkylidene; R3, R4 = H, alk(en/yn)yl, acyloxy, aryl, heterocyclyl, OH or NH2 or SH or derivs.; or R3R4 = O, S, NH, NOH, NNH2, CH2 or derivs.; R5, R6 = H, (un) substituted alk(en/yn)yl, cycloalkyl, aryl, heterocyclyl, OH or SH or derivs.; R7, R8 = H, alkyl] and their derivs. and analogs are provided. Also provided are pharmaceutical compns. containing I and methods of their medical use, particularly in the prophylaxis and treatment of hyperlipidemic conditions, such as those associated with atherosclerosis or hypercholesterolemia. For instance, the keto aldehyde II was cyclized by Zn/TiCl3, and the resultant cycloolefin was oxidized and epoxidized by m-ClC6H4C(O)OOH and hydrogenated over Pd/C to give epimeric title compds.  $\alpha$ - and  $\beta$ -III in 25% and 13% yield, plus addnl. compds. In a test for inhibition of IBAT-mediated uptake of [14C]-taurocholate in H14 cells in vitro,  $\beta$ -III had an IC50 of 5  $\mu$ M.

IC ICM C07D337-08

ICS C07D409-10; C08G065-329; A61K031-38

CC 27-21 (Heterocyclic Compounds (One Hetero Atom))

Section cross-reference(s): 1

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     (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT
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(preparation of benzothiepines as antihyperlipidemics)

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study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); USES (Uses)
   (preparation of benzothiepines as antihyperlipidemics)
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RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); USES (Uses)
   (preparation of benzothiepines as antihyperlipidemics)
197373-50-5P 197373-51-6P 197373-52-7P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological
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study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(preparation of benzothiepines as antihyperlipidemics)

RN197373-50-5 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-5-(3nitrophenyl) -, 1,1-dioxide, (4R,5R) -rel - (9CI) (CA INDEX NAME)

RN 197373-51-6 HCAPLUS

CN 1-Benzothiepin-4-ol, 5-(3-aminophenyl)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197373-52-7 HCAPLUS

CN Pentanamide, 5-bromo-N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

IT 197373-37-8P 197373-54-9P 197374-04-2P 197374-59-7P 197375-96-5P 197376-42-4P

19/3/4-59-7P 19/3/5-96-5P 19/3/6-42-197376-55-9P 197384-36-4P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of benzothiepines as antihyperlipidemics)

RN 197373-37-8 HCAPLUS

CN 1-Propanesulfonic acid, 3-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-, rel- (9CI) (CA INDEX NAME)

## Relative stereochemistry.

RN 197373-54-9 HCAPLUS

CN 1-Pentanaminium, 5-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-5-oxo-, rel-, salt with trifluoroacetic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 197373-53-8 CMF C37 H60 N3 O4 S

## Relative stereochemistry.

CM 2

CRN 14477-72-6 CMF C2 F3 O2

RN 197374-04-2 HCAPLUS

CN 2-Propenamide, N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197374-59-7 HCAPLUS

CN Propanamide, 3-bromo-N-[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]-, rel- (9CI) (CA INDEX NAME)

RN 197375-96-5 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-5-[2-(dimethylamino)phenyl]-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel-(9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 197376-42-4 HCAPLUS

CN Benzenaminium, 3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]-N,N,N-trimethyl-, iodide, rel-(9CI) (CA INDEX NAME)

Relative stereochemistry.

• I-

RN 197376-55-9 HCAPLUS

CN 1-Benzothiepin-4-ol, 3,3-dibutyl-7-(dimethylamino)-5-[3-(dimethylamino)phenyl]-2,3,4,5-tetrahydro-, 1,1-dioxide, (4R,5R)-rel-(9CI) (CA INDEX NAME)

RN 197384-36-4 HCAPLUS

CN 1-Hexanaminium, 6-[[3-[(4R,5R)-3,3-dibutyl-7-(dimethylamino)-2,3,4,5-tetrahydro-4-hydroxy-1,1-dioxido-1-benzothiepin-5-yl]phenyl]amino]-N,N,N-triethyl-6-oxo-, rel-, salt with trifluoroacetic acid (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 197384-35-3 CMF C38 H62 N3 O4 S

Relative stereochemistry.

CM 2

CRN 14477-72-6 CMF C2 F3 O2

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L26 ANSWER 1 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 1

ACCESSION NUMBER:

2005:144437 HCAPLUS

DOCUMENT NUMBER:

142:233695

TITLE:

Heterologous expression of human  $\alpha6\beta4\beta3\alpha5$  nicotinic acetylcholine

receptors: binding properties consistent with their natural expression require quaternary subunit assembly

including the  $\alpha$ 5 subunit

AUTHOR(S):

Grinevich, Vladimir P.; Letchworth, Sharon R.;

Lindenberger, Kari A.; Menager, Jean; Mary, Veronique; Sadieva, Khalima A.; Buhlman, Lori M.; Bohme, Georg

Andrees; Pradier, Laurent; Benavides,

Jesus; Lukas, Ronald J.; Bencherif, Merouane

CORPORATE SOURCE: Targacept, Inc., Winston-Salem, NC, USA

Journal of Pharmacology and Experimental Therapeutics SOURCE:

(2005), 312(2), 619-626

CODEN: JPETAB; ISSN: 0022-3565

PUBLISHER: American Society for Pharmacology and Experimental

Therapeutics

DOCUMENT TYPE: Journal English LANGUAGE:

Heterologous expression and lesioning studies were conducted to identify possible subunit assembly partners in nicotinic acetylcholine receptors (nAChR) containing α6 subunits (α6\* nAChR). SHEP1 human epithelial cells were transfected with the requisite subunits to achieve stable expression of human  $\alpha6\beta2$ ,  $\alpha6\beta4$ ,  $\alpha6\beta2\beta3$ ,  $\alpha6\beta4\beta3$ , or  $\alpha6\beta4\beta3\alpha5$  nAChR. Cells expressing subunits needed to form α6β4β3α5 nAChR exhibited saturable [3H] epibatidine binding (Kd = 95.9 pM and Bmax = 84.5 fmol/mg of protein). The rank order of binding competition potency (Ki) for prototypical nicotinic compds. was  $\alpha$ -conotoxin MII (6 nM) > nicotine (156 nM) .apprx. methyllycaconitine (200 nM) >  $\alpha$ -bungarotoxin (>10  $\mu$ M),

similar to that for nAChR in dopamine neurons displaying a distinctive

pharmacol. 6-Hydroxydopamine lesioning studies indicated that  $\beta 3$  and  $\alpha 5$  subunits are likely partners of the  $\alpha 6$  subunits in nAChR expressed in dopaminergic cell bodies. Similar to findings in rodents, quant. real-time reverse transcription-polymerase chain reactions of human brain indicated that  $\alpha 6$  subunit mRNA expression was 13-fold higher in the substantia nigra than in the cortex or the rest of the brain. Thus, heterologous expression studies suggest that the human  $\alpha 5$  subunit makes a critical contribution to  $\alpha 6\beta 4\beta 3\alpha 5$  nAChR assembly into a ligand-binding form with native  $\alpha 6*$ -nAChR-like pharmacol. and of potential physiol. and pathophysiol. relevance.

REFERENCE COUNT: 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 2 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 2

ACCESSION NUMBER: 2005:47732 HCAPLUS

DOCUMENT NUMBER: 143:76040

TITLE: ABCA2 is a strong genetic risk factor for early-onset

Alzheimer's disease

AUTHOR(S): Mace, Sandrine; Cousin, Emmanuelle; Ricard, Sylvain;

Genin, Emmanuelle; Spanakis, Emmanuel;

Lafargue-Soubigou, Carole; Genin, Berengere; Fournel, Raphael; Roche, Sandrine; Haussy, Gilles; Massey, Florence; Soubigou, Stephane; Brefort, Georges; Benoit, Patrick; Brice, Alexis; Campion, Dominique;

Hollis, Melvyn; Pradier, Laurent;

Benavides, Jesus; Deleuze, Jean-Francois Aventis Pharma, Evry Genetics Center and

Neurodegenerative Disease Group, Paris Research

Center, Vitry-sur-Seine, 94400, Fr.

Neurobiology of Disease (2005), 18(1), 119-125

CODEN: NUDIEM; ISSN: 0969-9961

PUBLISHER: Elsevier
DOCUMENT TYPE: Journal
LANGUAGE: English

CORPORATE SOURCE:

SOURCE:

Recent epidemiol., biol. and genetic data indicate a relationship between cholesterol and Alzheimer's disease (AD) including the association of polymorphisms of ABCA1 (a gene that is known to participate in cholesterol and phospholipid transport) with AD prevalence. Based on these data, we postulated that genetic variation in the related and brain-specific ABCA2 gene leads to increase risk of AD. A large case-control study was conducted where the sample was randomly divided into a hypothesis-testing sample (230 cases/286 controls) and a validation sample (210 cases/233 controls). Among the 45 SNPs we tested, one synonymous SNP (rs908832) was found significantly associated with AD in both samples. Addnl. analyses performed on the whole sample showed a very strong association between this marker and early-onset AD (OR = 3.82, 95% C.I. = [2.00 - 7.30], P = 5+ 10-5). Further research is needed to understand the functional role of this polymorphism. However, together with the reported assocns. of AD with APOE, CYP46Al and ABCAl, the present result adds a very significant support for the role of cholesterol and phospholipid homeostasis in AD and a rationale for testing novel cholesterol homeostasis-related therapeutic strategies in AD.

REFERENCE COUNT: 34 THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 3 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 3

ACCESSION NUMBER: 2005:978141 HCAPLUS

DOCUMENT NUMBER: 144:168853

TITLE: Unraveling substantia nigra sequential gene expression

in a progressive MPTP-lesioned macaque model of

Parkinson's disease

Bassilana, F.; Mace, N.; Li, Q.; Stutzmann, J. M.; AUTHOR (S):

Gross, C. E.; Pradier, L.; Benavides,

J.; Menager, J.; Bezard, E.

CORPORATE SOURCE:

Sanofi-Aventis, Vitry sur Seine, Fr.

SOURCE:

Neurobiology of Disease (2005), 20(1), 93-103

CODEN: NUDIEM; ISSN: 0969-9961

PUBLISHER:

Elsevier Journal

DOCUMENT TYPE: LANGUAGE:

English

Taking advantage of a progressive nonhuman primate model mimicking Parkinson's disease (PD) evolution, we monitored transcriptional fluctuations in the substantia nigra using Affymetrix microarrays in control (normal), saline-treated (normal), 6 days-treated (asymptomatic with 20% cell loss), 12 days-treated (asymptomatic with 40% cell loss) and 25 days-treated animals (fully parkinsonian with 85% cell loss). Two statistical methods were used to ascertain the regulation and real-time quant. PCR was used to confirm their regulation. Surprisingly, the number of deregulated transcripts is limited at all time points and five clusters exhibiting different profiles were defined using a hierarchical clustering algorithm. Such profiles are likely to represent activation/deactivation of mechanisms of different nature. We briefly speculate about (i) the existence of yet unknown compensatory mechanisms is unraveled, (ii) the putative triggering of a developmental program in the mature brain in reaction to progressing degeneration and finally, (iii) the activation of mechanisms leading eventually to death in final stage. These data should help development of new therapeutic approaches either aimed at enhancing existing compensatory mechanisms or at protecting dopamine neurons.

REFERENCE COUNT: THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS 35 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 4 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 4

ACCESSION NUMBER: 2004:1010181 HCAPLUS

DOCUMENT NUMBER: 142:21640

TITLE: Amyloid β-induced Changes in Nitric Oxide Production and Mitochondrial Activity Lead to

Apoptosis

AUTHOR (S): Keil, Uta; Bonert, Astrid; Marques, Celio A.;

Scherping, Isabel; Weyermann, Joerg; Strosznajder, Joanna B.; Mueller-Spahn, Franz; Haass, Christian;

Czech, Christian; Pradier, Laurent; Mueller,

Walter E.; Eckert, Anne

CORPORATE SOURCE: Departments of Pharmacology, Biocenter, University of

Frankfurt, Frankfurt, 60439, Germany

Journal of Biological Chemistry (2004), 279(48), SOURCE:

50310-50320

CODEN: JBCHA3; ISSN: 0021-9258

PUBLISHER: American Society for Biochemistry and Molecular

> Biology Journal

DOCUMENT TYPE: LANGUAGE:

English

Increasing evidence suggests an important role of mitochondrial dysfunction in the pathogenesis of Alzheimer's disease. Thus, the authors investigated the effects of acute and chronic exposure to increasing concns. of amyloid  $\beta$  (A $\beta$ ) on mitochondrial function and nitric oxide (NO) production in vitro and in vivo. The authors' data demonstrate that PC12 cells and human embryonic kidney cells bearing the Swedish double mutation in the amyloid precursor protein gene (APPsw), exhibiting

substantial Aß levels, have increased NO levels and reduced ATP levels. The inhibition of intracellular Aß production by a functional y- secretase inhibitor normalizes NO and ATP levels, indicating a direct involvement of AB in these processes. Extracellular treatment of PC12 cells with comparable AB concns. only leads to weak changes, demonstrating the important role of intracellular AB. In 3-mo-old APP transgenic (tg) mice, which exhibit no plagues but already detectable AB levels in the brain, reduced ATP levels can also be observed showing the in vivo relevance of the authors' findings. Moreover, the authors could demonstrate that APP is present in the mitochondria of APPsw PC12 cells. This presence might be directly involved in the impairment of cytochrome c oxidase activity and depletion of ATP levels in APPsw PC12 cells. In addition, APPsw human embryonic kidney cells, which produce 20-fold increased AB levels compared with APPsw PC12 cells, and APP to mice already show a significantly decreased mitochondrial membrane potential under basal conditions. The authors suggest a hypothetical sequence of pathogenic steps linking mutant APP expression and amyloid production with enhanced NO production and mitochondrial dysfunction finally leading to cell death.

REFERENCE COUNT:

THERE ARE 79 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 5 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 5

ACCESSION NUMBER:

2004:910579 HCAPLUS 142:4782

DOCUMENT NUMBER: TITLE:

Massive CA1/2 neuronal loss with intraneuronal and N-terminal truncated Aβ42 accumulation in a novel

Alzheimer transgenic model

AUTHOR (S):

Casas, Caty; Sergean, Nicolas; Itier, Jean-Michel; Blanchard, Veronique; Wirths, Oliver; van der Kolk, Nicolien: Vingtdeux, Valerie: van de Steeg, Evita;

Ret, Gwenaelle; Canton, Thierry; Drobecq,

Herve; Clark, Allan; Bonici, Bruno; Delacourte, Andre;

Benavides, Jesus; Schmitz, Christoph; Tremp, Gunter; Bayer, Thomas A.; Benoit, Patrick;

Pradier, Laurent

CORPORATE SOURCE:

Departments of Central Nervous System/Alzheimer Disease, INSERM U422, Aventis-Pharma Paris Research

Center, Paris, Fr.

SOURCE:

American Journal of Pathology (2004), 165(4),

1289-1300

CODEN: AJPAA4; ISSN: 0002-9440

PUBLISHER:

American Society for Investigative Pathology

DOCUMENT TYPE:

Journal LANGUAGE: English

Alzheimer's disease (AD) is characterized by a substantial degeneration of pyramidal neurons and the appearance of neuritic plaques and neurofibrillary tangles. Here we present a novel transgenic mouse model, APPSLPS1KI that closely mimics the development of AD-related neuropathol. features including a significant hippocampal neuronal loss. transgenic mouse model carries M233T/L235P knocked-in mutations in presenilin-1 and overexpresses mutated human β-amyloid (Aβ) precursor protein. A $\beta$ x-42 is the major form of A $\beta$  species present in this model with progressive development of a complex pattern of N-truncated variants and dimers, similar to those observed in AD brain. 10 mo of age, an extensive neuronal loss (>50%) is present in the CA1/2 hippocampal pyramidal cell layer that correlates with strong accumulation of intraneuronal Aβ and thioflavine-S-pos. intracellular material but

not with extracellular Aβ deposits. A strong reactive astrogliosis develops together with the neuronal loss. This loss is already detectable at 6 mo of age and is PSIK1 gene dosage-dependent. Thus, APPSLPS1KI mice. further confirm the critical role of intraneuronal  $A\beta42$  in neuronal loss and provide an excellent tool to investigate therapeutic strategies designed to prevent AD neurodegeneration.

REFERENCE COUNT:

THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS 48 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 6 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 6

ACCESSION NUMBER:

2004:984714 HCAPLUS

DOCUMENT NUMBER:

142:169848

TITLE:

In vitro and in vivo characterization of TC-1827, a

novel brain  $\alpha 4\beta 2$  nicotinic receptor agonist

with pro-cognitive activity

AUTHOR (S):

Bohme, Georg Andrees; Letchworth, Sharon R.; Piot-Grosjean, Odile; Gatto, Gregory J.; Obinu, Marie-Carmen; Caldwell, William S.; Laville, Michel;

Brunel, Pascale; Pellerin, Rachel; Leconte, Jean-Pierre; Genevois-Borella, Arielle; Dubedat, Pierre; Mazadier, Martine; Pradier, Laurent;

Bencherif, Merouane; Benavides, Jesus

CORPORATE SOURCE:

Centre de Recherches de Paris, Aventis Pharma S.A.,

Vitry Sur Seine, Fr.

SOURCE:

Drug Development Research (2004), 62(1), 26-40

CODEN: DDREDK; ISSN: 0272-4391

PUBLISHER: Wiley-Liss, Inc.

DOCUMENT TYPE:

Journal

42

LANGUAGE: English

Nicotine activates specific receptors that are cation-permeable ionic channels located in the central and autonomous nervous systems, as well as at the neuromuscular junction. Administration of nicotine to animals and humans has been shown to enhance cognitive processes. However, side effects linked to the activation of peripheral nicotinic receptors limit the usefulness of nicotine for the treatment of cognitive disorders such as Alzheimer's disease (AD) or mild cognitive impairments (MCI). The synthesis and properties of TC-1827, a novel metanicotine derivative that activates brain  $\alpha4\beta2$  nicotinic receptors is described. TC-1827 has high affinity for nicotine-labeled receptors in the cortex (Ki = 34 nM), full-agonist intrinsic activity in  $\alpha 4\beta 2$ -mediated neurotransmitter release studies in synaptosomes, and has no functional activity at nicotinic receptors in ganglionic or muscular cell lines. The compound enhances long-term potentiation in hippocampal slices, a form of synaptic plasticity thought to be involved in information storage at the cellular level. In vivo studies demonstrate that TC-1827 dose-dependently occupies thalamic nicotinic receptors labeled with [3H]-cytisine, increases cortical extracellular acetylcholine levels following oral administration, and enhances cognitive performance in rat and mice behavioral procedures of learning and memory. Pharmacokinetic studies in mice, rats, and monkeys indicated that TC-1827 has good oral absorption with a first pass effect resulting in bioavailabilities of 13-65% across dose/species. Cardiovascular safety studies indicate good cardiovascular tolerability for this compound. The present data demonstrate that TC-1827 is a selective and potent activator of brain  $\alpha 4\beta 2$  nicotinic receptors and is a prototypical member of a new class of compds. with potential utility in the symptomatic treatment of cognitive disorders including AD and MCI.

REFERENCE COUNT:

THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L26 ANSWER 7 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 7

ACCESSION NUMBER: 2003:704890 HCAPLUS

DOCUMENT NUMBER: 139:321545

TITLE: Parkin gene inactivation alters behaviour and dopamine

neurotransmission in the mouse

AUTHOR(S): Itier, Jean-Michel; Ibanez, Pablo; Mena, Maria

Angeles; Abbas, Nacer; Cohen-Salmon, Charles; Bohme, Georg Andrees; Laville, Michel; Pratt, Jeremy; Corti,

Olga; Pradier, Laurent; Ret, Gwenaeelle;

Joubert, Chantal; Periquet, Magali; Araujo, Francisco;

Negroni, Julia; Casarejos, Maria Jose; Canals, Santiago; Solano, Rosa; Serrano, Alba; Gallego, Eva;

Sanchez, Marina; Denefle, Patrice; Benavides, Jesus; Tremp, Guenter; Rooney, Thomas A.; Brice,

Alexis; Garcia de Yebenes, Justo

CORPORATE SOURCE: Functional Genomics Department, Aventis Pharma SA,

Vitry-sur-Seine, F-94400, Fr.

SOURCE: Human Molecular Genetics (2003), 12(18), 2277-2291

CODEN: HMGEE5: ISSN: 0964-6906

PUBLISHER: Oxford University Press

DOCUMENT TYPE: Journal LANGUAGE: English

Mutations of the parkin gene are the most frequent cause of early onset autosomal recessive parkinsonism (EO-AR). Here the authors show that inactivation of the parkin gene in mice results in motor and cognitive deficits, inhibition of amphetamine-induced dopamine release and inhibition of glutamate neurotransmission. The levels of dopamine are increased in the limbic brain areas of parkin mutant mice and there is a shift towards increased metabolism of dopamine by MAO. Although there was no evidence for a reduction of nigrostriatal dopamine neurons in the parkin mutant mice, the level of dopamine transporter protein was reduced in these animals, suggesting a decreased d. of dopamine terminals, or adaptative changes in the nigrostriatal dopamine system. GSH levels were increased in the striatum and fetal mesencephalic neurons from parkin mutant mice, suggesting that a compensatory mechanism may protect dopamine neurons from neuronal death. These parkin mutant mice provide a valuable tool to better understand the preclin. deficits observed in patients with PD and to characterize the mechanisms leading to the degeneration of dopamine neurons that could provide new strategies for neuroprotection.

REFERENCE COUNT: 91 THERE ARE 91 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 8 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 8

ACCESSION NUMBER:

2003:235026 HCAPLUS

DOCUMENT NUMBER:

139:143701

TITLE:

Synthesis of a biotin-tagged photoaffinity probe of

2-azetidinone cholesterol absorption

inhibitors

AUTHOR(S):

Frick, Wendelin; Bauer-Schafer, Andrea; Bauer, Jochen;

Girbig, Frank; Corsiero, Daniel; Heuer, Hubert

; Kramer, Werner

CORPORATE SOURCE:

Disease Group Metabolic Diseases Industriepark Hochst, Aventis Pharma Deutschland GmbH, Frankfurt am Main,

D-65926, Germany

SOURCE:

Bioorganic & Medicinal Chemistry (2003), 11(8),

1639-1642

CODEN: BMECEP; ISSN: 0968-0896

PUBLISHER:

Elsevier Science Ltd.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

The design and synthesis of a biotin-tagged photoreactive analog C-4 of

the cholesterol absorption inhibitor Ezetimibe is

described. Photoaffinity labeling of intestinal brush border

membrane vesicles with C-4 and subsequent streptavidin-biotin chromatog. leads to selective extraction of a 145 kDa integral membrane protein as the

mol. target for cholesterol absorption inhibitors.

REFERENCE COUNT:

THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS 10

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 9 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 9

ACCESSION NUMBER:

2003:903740 HCAPLUS

DOCUMENT NUMBER:

140:179480

TITLE:

Time sequence of maturation of dystrophic neurites

associated with Aß deposits in APP/PS1 transgenic

mice

AUTHOR (S):

Blanchard, Veronique; Moussaoui, Saliha; Czech,

Christian; Touchet, Nathalie; Bonici, Bruno; Planche,

Michel; Canton, Thierry; Jedidi, Iness;

Gohin, Micheline; Wirths, Oliver; Bayer, Thomas A.; Langui, Dominique; Duyckaerts, Charles; Tremp, Gunter;

Pradier, Laurent

CORPORATE SOURCE:

Neurodegenerative Disease Group, Centre de Recherche

de Paris, Vitry sur Seine, 94403, Fr.

SOURCE:

Experimental Neurology (2003), 184(1), 247-263

CODEN: EXNEAC; ISSN: 0014-4886

PUBLISHER:

Elsevier Science

DOCUMENT TYPE:

Journal English

LANGUAGE:

Several novel transgenic mouse models expressing different mutant APPs in combination with mutant PS1 have been developed. These models have been analyzed to investigate the formation and progressive alterations of dystrophic neurites (DNs) in relation to Aβ deposits. In the most aggressive model,  $A\beta$  deposits appear as early as 2.5 mo of age. Maturation of DNs was qual. quite similar among models and in some respect reminiscent of human AD pathol. From the onset of deposition, most if not all Aβ deposits were decorated with a high number of APP-, ubiquitin-, and MnSOD-immunoreactive DNs. Phosphorylated Tau DNs, however, appeared at a much slower rate and were more restricted. Mitochondrial dysfunction markers were observed in DNs: the frequency and the d. per deposit of DNs accumulating cytochrome c, cytochrome oxidase 1, and Bax progressively increased with age. Later, the burden of reactive DNs was reduced around large compact/mature deposits. In addition, the previously described phenomenon of early intraneuronal AB accumulation in our models was associated with altered expression of APP protein as well as oxidative and mitochondrial stress markers occasionally in individual neurons. The present study demonstrates that oxidative and mitochondrial stress factors are present at several phases of AB pathol. progression, confirming

REFERENCE COUNT:

the neuronal dysfunction in APP transgenic mice. 75 THERE ARE 75 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 10 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 10

139:83282

ACCESSION NUMBER:

2003:327208 HCAPLUS

DOCUMENT NUMBER: TITLE:

A risk for early-onset Alzheimer's disease associated

AUTHOR(S):

with the APBB1 gene (FE65) intron 13 polymorphism Cousin, Emmanuelle; Hannequin, Didier; Ricard,

Sylvain; Mace, Sandrine; Genin, Emmanuelle; Chansac,

Celine; Brice, Alexis; Dubois, Bruno; Frebourg,

Thierry; Mercken, Luc; Benavides, Jesus;

Pradier, Laurent; Campion, Dominique; Deleuze,

Jean-Francois

CORPORATE SOURCE: Paris Research Center, Evry Genetics Center &

Neurodegenerative Disease Group, Aventis Pharma,

Vitry-sur-Seine, 94400, Fr.

Neuroscience Letters (2003), 342(1,2), 5-8 SOURCE:

CODEN: NELED5; ISSN: 0304-3940

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

Alzheimer's disease (AD) is a genetically complex neurodegenerative disorder and the leading cause of dementia of the elderly. Recently, Hu et al. suggested that a trinucleotide deletion in intron 13 of the APBB1 gene was a factor protecting against late-onset AD. We report here the results of a case/control study aimed at replicating this association Our study included 461 AD patients and 397 matched controls. We compared the allele and genotype frequencies of the polymorphism between the two groups but did not find any statistically significant difference (P=0.08 and P=0.09, resp.). By contrast, adjusting for age and sex, we found a slight risk associated with the deletion (odds ratio=1.47, 95% confidence interval=1.05-2.04). Stratification by age showed that the risk effect associated with the deletion concerned subjects aged less than 65 yr.

THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 14 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 11 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 11

ACCESSION NUMBER: 2002:471429 HCAPLUS

DOCUMENT NUMBER: 138:71094

Amyloid precursor protein family-induced neuronal TITLE:

death is mediated by impairment of the neuroprotective

calcium/calmodulin protein kinase IV-dependent

signaling pathway

Mbebi, Corinne; See, Violaine; Mercken, Luc; AUTHOR (S):

Pradier, Laurent; Muller, Ulrike; Loeffler,

Jean-Philippe

CORPORATE SOURCE: Universite Louis Pasteur, Faculte de Medecine, EA 3433

Molecular signaling and neurodegeneration, Strasbourg,

67000, Fr.

Journal of Biological Chemistry (2002), 277(23), SOURCE:

20979-20990

CODEN: JBCHA3; ISSN: 0021-9258

American Society for Biochemistry and Molecular PUBLISHER:

Biology

DOCUMENT TYPE: Journal

LANGUAGE: English

The aberrant metabolism of  $\beta$ -amyloid precursor protein (APP) and the progressive deposition of its derived fragment β-amyloid peptide are early and constant pathol. hallmarks of Alzheimer's disease. Because APP is able to function as a cell surface receptor, the authors investigated here whether a disruption of the normal function of APP may contribute to the pathogenic mechanisms in Alzheimer's disease. To this aim, the authors generated a specific chicken polyclonal antibody directed against the extracellular domain of APP, which is common with the  $\beta$ -amyloid precursor-like protein type 2. Exposure of cultured cortical neurons to this antibody (APP-Ab) induced

cell death preceded by neurite degeneration, oxidative stress, and nuclear

condensation. Interestingly, caspase-3-like protease was not activated in this neurotoxic action suggesting a different mode of cell death than classical apoptosis. Further anal. of the mol. mechanisms revealed a calpain- and calcineurin-dependent proteolysis of the neuroprotective calcium/calmodulin-dependent protein kinase IV and its nuclear target protein cAMP responsive element binding protein. These effects were abolished by the G protein inhibitor pertussis toxin, strongly suggesting that APP binding operates via a GTPase-dependent pathway to cause neuronal death.

REFERENCE COUNT:

90 THERE ARE 90 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 12 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 12

ACCESSION NUMBER:

2002:81635 HCAPLUS

DOCUMENT NUMBER:

136:307775

TITLE:

Increased Expression of Presenilin 2 Inhibits

Protein Synthesis

AUTHOR (S):

Gamliel, Amir; Teicher, Carmit; Michaelson, Daniel'M.;

Pradier, Laurent; Hartmann, Tobias; Beyreuther, Konrad; Stein, Reuven

CORPORATE SOURCE:

Department of Neurobiochemistry, George S. Wise Faculty of Life Sciences, Tel Aviv University, Tel

Aviv-Jaffa, 69978, Israel

SOURCE:

Molecular and Cellular Neuroscience (2002), 19(1),

111-124

CODEN: MOCNED; ISSN: 1044-7431

PUBLISHER:

Academic Press

DOCUMENT TYPE:

Journal English

LANGUAGE: Mutations in the presentlin genes PS1 and PS2 are a major cause of early onset familial Alzheimer's disease (AD). Previous studies have suggested that presentlins have several functions, including  $\gamma$ secretase activity. It was also shown that presentlin expression is increased in the brains of some AD patients and ischemic rodents. present study examines the effect of increased presenilin expression on protein synthesis. We show here that overexpression of wild-type PS2 (PS2wt) or PS2 mutant containing the FAD mutation N141I (PS2mut) in various cell lines inhibits the synthesis of coexpressed reporter and endogenous proteins. Furthermore, endogenous PS2 seems to be needed for translation inhibition since PS2 null fibroblasts were translationally more active than PS2+/+ fibroblasts under conditions known to inhibit translation. Overexpression of PS1 also appeared to cause inhibition of protein synthesis, but its effect was much weaker than that of PS2. Taken together, the results suggest that increased expression of PS2 and possibly also of PS1 inhibits translation and that presenilins may function as regulators of protein synthesis. (c) 2002 Academic Press.

REFERENCE COUNT:

67 THERE ARE 67 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 13 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 13

ACCESSION NUMBER:

2001:24031 HCAPLUS

DOCUMENT NUMBER:

134:176079

TITLE:

Identification of binding proteins for cholesterol absorption inhibitors as

components of the intestinal

cholesterol transporter

AUTHOR(S):

Kramer, W.; Glombik, H.; Petry, S.; Heuer, H.

; Schafer, H.-L.; Wendler, W.; Corsiero, D.; Girbig,

F.; Weyland, C.

CORPORATE SOURCE:

Disease Group Metabolic Diseases, Aventis Pharma

Deutschland GmbH, Frankfurt am Main, D-65926, Germany

SOURCE: FEBS Letters (2000), 487(2), 293-297

CODEN: FEBLAL; ISSN: 0014-5793

PUBLISHER:

Elsevier Science B.V.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

To identify protein components of the intestinal

cholesterol transporter, rabbit small intestinal brush

border membrane vesicles were submitted to photoaffinity labeling using photoreactive derivs. of 2-azetidinone cholesterol absorption inhibitors. An integral membrane protein of Mr 145.3±7.5 kDa

was specifically labeled in brush border membrane vesicles from rabbit jejunum and ileum. Its labeling was concentration-dependently inhibited

by the presence of cholesterol absorption inhibitors

whereas bile acids, D-glucose, fatty acids or cephalexin had no effect.

The inhibitory potency of 2-azetidinones to inhibit

photolabeling of the 145 kDa protein correlated with their in vivo activity to inhibit intestinal cholesterol

absorption. These results suggest that an integral membrane protein of Mr 145 kDa is (a component of) the cholesterol absorption system in

REFERENCE COUNT:

THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS 34

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 14 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 14

the brush border membrane of small intestinal enterocytes.

ACCESSION NUMBER:

1999:807208 HCAPLUS

DOCUMENT NUMBER:

132:146489

TITLE:

Alterations of carbohydrate and lipid intermediary

metabolism during inhibition of glucose-6-phosphatase in rats

AUTHOR (S):

Herling, A. W.; Burger, H.-J.; Schubert, G.; Hemmerle,

H.; Schaefer, H.-L.; Kramer, W.

CORPORATE SOURCE:

H 821 Pharmacology, Hoechst Marion Roussel Deutschland

GmbH, Frankfurt am Main, 65926, Germany

SOURCE:

European Journal of Pharmacology (1999), 386(1), 75-82

CODEN: EJPHAZ; ISSN: 0014-2999

PUBLISHER:

Elsevier Science B.V.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

S 4048 (1-[2-(4-Chloro-phenyl)-cyclopropylmethoxy]-3,4-dihydroxy-5-(3imidazo[4,5-b]pyridin-1-yl-3-phenyl-acryloyloxy)-cyclohexanecarboxylic acid), a derivative of chlorogenic acid, specifically inhibits the qlucose-6-phosphate translocating component T1 of the glucose-6phosphatase system. Its pharmacol. effect was studied on carbohydrate and lipid parameters in rats. In starved and fed rats, S 4048 caused a dose-dependent reduction of blood glucose levels with a corresponding increase in hepatic and renal glycogen and glucose-6-phosphate. The major quant. route of carbon flux in the liver during S 4048-induced inhibition of the glucose-6-phosphatase activity seemed to be glycogenesis. Plasma free fatty acids were increased secondarily due to the S 4048-induced hypoglycemia. Hepatic triglycerides were increased possibly due to increased re-esterification of the readily available free fatty acids. Glucose-6-phosphate translocase inhibitors may be useful for exptl. studying aspects of type 1 glycogen storage disease in laboratory

as well as for the therapeutic modulation of inappropriately high rates of hepatic glucose production in type 2 diabetes.

Krishnan 10/734,787 09/05/2006

REFERENCE COUNT:

38 THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 15 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 15

ACCESSION NUMBER:

1999:178553 HCAPLUS

DOCUMENT NUMBER:

131:3766

TITLE:

Mapping the APP/presenilin (PS) binding domains: the hydrophilic N-terminus of PS2 is sufficient for interaction with APP and can

displace APP/PS1 interaction

AUTHOR (S):

Pradier, Laurent; Carpentier, Nathalie; Delalonde, Laurence; Clavel, Nicole; Bock,

Marie-Dominique; Buee, Luc; Mercken, Luc; Tocque,

Bruno; Czech, Christian

CORPORATE SOURCE:

Gene Medicine Department, Rhone-Poulenc Rorer, Vitry,

94400, Fr.

SOURCE:

Neurobiology of Disease (1999), 6(1), 43-55

CODEN: NUDIEM; ISSN: 0969-9961

PUBLISHER:

Academic Press

Journal DOCUMENT TYPE: English LANGUAGE:

Mutations in presenilin 1 and presenilin 2 (PS1 and PS2, resp.) genes cause the large majority of familial forms of early-onset Alzheimer's disease. The phys. interaction between presentlins and APP has been recently described using coimmunopptn. With a similar technique, we confirmed this interaction and have mapped the interaction domains on both PS2 and APP. Using several carboxy-terminal truncated forms of PS2, we demonstrated that the hydrophilic amino terminus of PS2 (residues 1 to 87, PS2NT) was sufficient for interaction with APP. Interestingly, only a construct with a leader peptide for secretion (SecPS2NT) and not its cytosolic counterpart was shown to interact with APP. For APP, we could demonstrate interaction of PS2 with the last 100 but not the last 45 amino acids of APP, including therefore the AB region. Accordingly, SecPS2NT is capable of binding to AB-immunoreactive species in conditioned medium. In addition, a second region in the extracellular domain of APP also interacted with PS2. Comparable results with PS1 indicate that the two presenilins share similar determinants of binding to APP. Confirming these results, SecPS2NT is able to inhibit PS1/ APP interaction. Such a competition makes it unlikely that the PS/APP interaction results from nonspecific aggregation of PS in transfected cells. The phys. interaction of presenilins with a region encompassing the Aß sequence of APP could be causally related to the misprocessing of APP and the production of (c) 1999 Academic Press. Aβ1-42.

REFERENCE COUNT:

43 THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 16 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 16

ACCESSION NUMBER:

1996:612230 HCAPLUS

DOCUMENT NUMBER:

125:293960

TITLE:

Molecular cloning, functional expression,

pharmacological characterization and chromosomal localization of the human metabotropic glutamate

receptor type 3

AUTHOR (S):

Emile, Lydia; Mercken, Luc; Apiou, Francoise;

Pradier, Laurent; Bock, Marie-Dominique;

Menager, Jean; Clot, Josette; Doble, Adam; Blanchard,

Jean-Charles

CORPORATE SOURCE: Rhone-Poulenc Rorer SA, Centre de Recherche de

Vitry-Alfortville, Vitry sur Seine, 94400, Fr.

SOURCE: Neuropharmacology (1996), 35(5), 523-530

CODEN: NEPHBW; ISSN: 0028-3908

PUBLISHER: Elsevier
DOCUMENT TYPE: Journal
LANGUAGE: English

Glutamic acid is the major excitatory amino acid of the central nervous system which interacts with two receptor families, the ionotropic and metabotropic glutamate receptors. The metabotropic glutamate receptors (mGluRs) are coupled to G proteins and can be divided into three subgroups based on their sequence homol., signal transduction pathway and pharmacol. In this study, we describe the cloning of the cDNA encoding the human metabotropic glutamate receptor type 3 (HmGluR3). It was obtained by reverse transcription-polymerase chain reaction (RT-PCR) with degenerate oligonucleotides corresponding to highly conserved sequences between rat mGluRs. The receptor shows 879 amino acids with 96% amino acid sequence identity with rat mGluR3. It is strongly expressed in fetal and adult whole brain, especially in caudate nucleus and corpus callosum. gene was identified by fluorescence in situ hybridization on chromosome 7 band q22. Activation of the human mGluR3, permanently expressed in Baby Hamster Kidney (BHK) cells, by excitatory amino acids inhibits the forskolin-stimulated accumulation of intracellular cAMP. The rank order of potency is L-glutamic acid>> (1S,3R)-1-aminocyclopentane-1,3dicarboxylic acid ((1S, 3R) - ACPD) »ibotenic acid >quisqualic acid. (RS)- $\alpha$ -methyl-4-carboxyphenylglycine [(RS)-MCPG, 1 mM] is without effect on inhibition of forskolin-induced cAMP accumulation by L-glutamic acid.

L26 ANSWER 17 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 17

ACCESSION NUMBER: 1993:420435 HCAPLUS

DOCUMENT NUMBER: 119:20435

TITLE: Inhibition by riluzole of electrophysiological

responses mediated by rat kainate and NMDA receptors

expressed in Xenopus oocytes

AUTHOR(S): Debono, Marc Williams; Le Guern, Joelle; Canton,

Thierry; Doble, Adam; Pradier, Laurent

Third P. Doron Program Program

CORPORATE SOURCE: Dep. Biol., Rhone-Poulenc Rorer S. A., Vitry sur

Seine, 94403, Fr.

SOURCE: European Journal of Pharmacology (1993), 235(2-3),

283-9

CODEN: EJPHAZ; ISSN: 0014-2999

DOCUMENT TYPE: Journal LANGUAGE: English

AB The effects of riluzole, an anticonvulsant and neuroprotective compound, on excitatory amino acid-evoked currents were studied in Xenopus laevis oocytes injected with mRNA from rat whole brain or cortex. Responses to kainic acid were blocked by riluzole (IC50 = 167 μM) as well as by the quinoxalinedione antagonists 6-cyano-7-nitroquinoxaline-2,3-dione (CNQX:IC50 = 0.21 μM) and 2,3-dihydroxy-6-nitro-7-sulfamoylbenzo[f]quinoxaline (NBQX: IC50 = 0.043 μM). NMDA receptor antagonist 2-amino-phosphonovaleric acid 92-APV) yielded an IC50 of 6.1 μM in this system. The inhibition by both riluzole and 2-APV was reversible and did not appear to be use dependent, unlike that of the channel blocker MK-801 ([+]-5-methyl-10,11-dihydro-5H-dibenzo[a,d]cyclohepten-1,10-imine maleate). It was impossible to demonstrate an interaction of riluzole with any of the known ligand sites on either the kainate or the radioligand binding studies. These results suggest a direct but non-competitive action of riluzole on ionotropic

glutamate receptors.

L26 ANSWER 18 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 18

ACCESSION NUMBER: 1990:456670 HCAPLUS

DOCUMENT NUMBER: 113:56670

TITLE: Tumor necrosis factor (TNF) and endotoxin prime

effects of PAF in vivo

Heuer, H. O.; Letts, G.; Meade, C. J. AUTHOR(S):

CORPORATE SOURCE: Dep. Pharmacol., Boehringer Ingelheim, Ingelheim,

D-6507, Germany

SOURCE: Journal of Lipid Mediators (1990), 2(Suppl.),

S101-S108

CODEN: JLMEEG; ISSN: 0921-8319

DOCUMENT TYPE: Journal LANGUAGE: English

The purpose of the present study in NMRI mice was to investigate the action of platelet-activating factor (PAF) on mortality and intestinal transit velocity, the interaction of endotoxin or tumor necrosis factor (TNF) with the effect of PAF on these parameters and the effect of the PAF antagonist WEB 2086 on the endotoxin/TNF- and PAF-induced changes. PAF at a high dose (200 µg/kg i.v.) increased mortality and reduced transit velocity. This effect was inhibited by WEB 2086 (0.01-0.5 mg/kg i.p.) in a dose-dependent manner. Pretreatment with endotoxin (10 µg/kg i.v.) or TNF (40 µg/kg i.v.) enhanced the activity of PAF resulting in increased mortality and reduced transit velocity. This enhanced activity of PAF in the case of pretreatment with endotoxin or TNF occurred at doses at which PAF, endotoxin or TNF given alone did not affect these parameters. The ability of endotoxin or TNF to enhance the effect of PAF was maximal if the time delay between endotoxin and subsequent PAF administration was about 1-2 h. WEB 2086 (0.01-1 mg/kg i.p.) inhibited this priming in a dose-dependent fashion. These findings support suggestions of a role for PAF in endotoxin shock and TNF-associated shock-like syndrome.

L26 ANSWER 19 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 19

1989:508737 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 111:108737

TITLE: Effects of a new and specific PAF-antagonist, WEB 2086, on PAF and endotoxin/tumor necrosis factor

induced changes in mortality and intestinal

transit velocity

AUTHOR(S): Heuer, Hubert

CORPORATE SOURCE: Dep. Pharmacol., Boehringer Ingelheim K.-G.,

Ingelheim, D-6507, Fed. Rep. Ger.

SOURCE: Progress in Clinical and Biological Research (1989),

308 (Vienna Shock Forum, 2nd, 1988), 919-24

CODEN: PCBRD2; ISSN: 0361-7742

DOCUMENT TYPE: Journal LANGUAGE: English

Endotoxin at ≤40 µg/kg and platelet-activating factor (PAF) at ≤100 µg/kg did not alter the gastrointestinal transit time or prove lethal to young mice. Single doses of tumor necrosis factor (TNF) also had no effects at 0.5 or 1.0 mg/kg, i.v. Pretreatment with TNF or endotoxin increased the lethal effects of PAF. Pretreatment with WEB 2086 at 0.1-1.0 mg/kg, i.p., 15 min prior to PAF inhibited the intestinal transit velocity changes from PAF plus TNF or endotoxin. WEB 2086 at 0.01-1.0 mg/kg, i.p., given 15 min before, inhibited the lethal synergism between TNF and subsequent PAF.

L26 ANSWER 20 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 20

ACCESSION NUMBER: 1987:531594 HCAPLUS

DOCUMENT NUMBER: 107:131594

TITLE: Circadian rhythm in the membrane of circulating human

blood cells: microviscosity and number of

benzodiazepine binding sites. A search for regulation

by plasma ions, nucleosides, proteins or hormones

AUTHOR(S): Levi, Francis; Benavides, Jesus; Touitou,

Yvan; Quarteronet, Dominique; Canton, Thierry

; Uzan, Andre; Auzeby, Andre; Gueremy, Claude; Sulon,

Jose; et al.

CORPORATE SOURCE: Fond. Adolphe de Rothschild, Paris, 75940, Fr.

SOURCE: Chronobiology International (1987), 4(2), 235-43

CODEN: CHBIE4; ISSN: 0742-0528

DOCUMENT TYPE: Journal LANGUAGE: English

Circadian rhythms in both the number of peripheral type binding sites for benzodiazepines in platelet membranes and the microviscosity of the erythrocyte membrane were demonstrated in healthy men. Neither variable appeared to be linked to each other, or be regulated by the plasma concns. of total or free cortisol, testosterone, K+, Mq2+, Ca2+, cAMP, cGMP, or proteins or by the erythrocyte concentration of Mq2+ or K+ or by the plasma cAMP:cGMP ratio or by the ratio of intra-erythrocyte:plasma concns. of Mg2+ or K+. A highly significant neg. correlation was found between the microviscosity of the erythrocyte membrane and the activity of the membrane-bound enzyme methyltransferase I. Such a correlation was validated both on raw data and on 24 h-means. A circadian rhythm in the activity of this enzyme was also demonstrated. Moreover, a highly significant correlation was also found between plasma transcortin concentration (TRC) and microviscosity, and between TRC and methyltransferase I activity. Such findings may constitute clues towards the understanding of the regulation of the circadian rhythm in the fluidity of the red blood cell membrane in man and quide future steps with regard to the role of this rhythm upon the availability of drug binding sites at the cell surface.

L26 ANSWER 21 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 21

ACCESSION NUMBER: 1986:546030 HCAPLUS

DOCUMENT NUMBER: 105:146030

TITLE: Circadian rhythm in peripheral type benzodiazepine

binding sites in human platelets

AUTHOR(S): Levi, Francis; Benavides, Jesus; Touitou,

Yvan; Quarterronet, Dominique; Canton, Thierry; Uzan, Andre; Gueremy, Claude; Le Fur, Gerard;

Reinberg, Alain

CORPORATE SOURCE: Fond. Adolphe de Rothschild, Paris, 75940, Fr.

SOURCE: Biochemical Pharmacology (1986), 35(15), 2623-5

CODEN: BCPCA6; ISSN: 0006-2952

DOCUMENT TYPE: Journal LANGUAGE: English

AB Peripheral benzodiazepine (I) receptors were found in platelet membranes from all human subjects examined and at all time points. Such sites demonstrated a high affinity (KD) for [3H]PK 11195 (a selective ligand for peripheral I binding sites) and a high capacity (Bmax). Nonetheless large interindividual differences in the 24-h mean values of Bmax, KD, and platelet count were observed A circadian rhythm was found and validated for both Bmax and platelet count, but not for KD. The maximum binding occurred at 0350 h; the difference between maximum values was .apprx.20% of the 24-h mean. Circadian changes in the binding capacity were not related to

either platelet count (differences in peak times of resp. circadian rhythms or the affinity (no circadian rhythm in KD). As a result, the circadian in the binding capacity most likely reflects that in the number of binding sites per platelet. Apparently, the expression of I binding sites in human platelet exhibits a circadian rhythm.

L26 ANSWER 22 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 22

ACCESSION NUMBER:

1986:123089 HCAPLUS

DOCUMENT NUMBER:

104:123089

TITLE:

Biochemical characterization and quantitative autoradiographic study of the binding sites for

indalpine, a 5-HT reuptake inhibitor

, in cat brain

AUTHOR (S):

Benavides, J.; Malgouris, C.; Daniel, M.; Savaki, H.; Uzan, A.; Gueremy, C.; Le Fur, G.

CORPORATE SOURCE: SOURCE:

Coll. France, Paris, 75231/05, Fr. Encephale (1985), 11(6), 247-54 CODEN: ENCEAN; ISSN: 0013-7006

DOCUMENT TYPE:

Journal

LANGUAGE:

French

Autoradiog. and biochem. (competitive displacement) studies with cat brain slices showed that 3H-labeled indalpine [63758-79-2] bound with high affinity to serotoninergic receptors, as shown previously for the rat brain. The total number of binding sites was 146 fmol/mg protein, and the affinity constant was 2.6 mM. These sites could be differentiated from [3H]imipramine binding sites by their requirement for Na+ and their competitive inhibition by serotonin. Color-coded images of the distribution of the binding sites in various areas of the cat brain are presented. This distribution suggested that indalpine would have a stronger effect on the limbic system than on the extrapyramidal system, which is consistent with its known antidepressant properties in humans.

L26 ANSWER 23 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2006:681037 HCAPLUS

DOCUMENT NUMBER:

145:145534

TITLE:

Preparation of sulfonyl pyrrolidines, and their use for increasing blood high-density lipoprotein level for treating dyslipidemia, diabetes and related

diseases

INVENTOR(S):

Keil, Stefanie; Schaefer, Hans-Ludwig;

Glien, Maike; Guessregen, Stefan; Wendler, Wolfgang;

Esswein, Marion

PATENT ASSIGNEE(S):

Sanofi-Aventis Deutschland GmbH, Germany

SOURCE:

PCT Int. Appl., 169 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent German

LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT	NO.			KIN	D :	DATE			APPL	ICAT:	ION I	NO.		D	ATE	
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WO 2006	0723	93		A2		2006	0713	,	WO 2	005-	EP13	772		2	0051	221
W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KM,	KN,	KP,	KR,
	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,
	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,
	SG,	SK,	SL,	SM,	SY,	TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,

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VN, YU, ZA, ZM, ZW

RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ,
CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH,
GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
KG, KZ, MD, RU, TJ, TM

PRIORITY APPLN. INFO.:

DE 2005-102005000666A 20050104
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\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

The invention is related to substituted sulfonyl pyrrolidines, including compds. of formula I [R1 = fluoro/alkyl, (un)substituted Ph, heterocyclyl, etc.; R2 = alkyl, (un)substituted Ph, heterocyclyl, etc.; R3-R5 = independently H, F, Cl, Br, NO2, alkyl, Ph, etc.; with the exclusion of certain compds.], and their physiol. acceptable salts, and their use as drugs for increasing blood HDL level. E.g., a multi-step synthesis starting from 4-methylbenzaldehyde, was given for pyrrolidine trans-II•TFA. I increased ATP-binding cassette protein A1 (ABCA1) expression and thereby increased production of cholesterol in blood HDL. I are useful for treating dyslipidemia, coronary circulation diseases, arteriosclerosis, diabetes, and metabolic syndrome.

L26 ANSWER 24 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:857559 HCAPLUS

DOCUMENT NUMBER: 141:314568

TITLE: Novel diphenyl azetidinone with improved physiological

characteristics, corresponding production method, medicaments containing said compound and use of the

latter

INVENTOR(S): Jaehne, Gerhard; Frick, Wendelin; Lindenschmidt,

Andreas; Flohr, Stefanie; Heuer, Hubert; Schaefer, Hans-Ludwig; Kramer, Werner; Galia,

Eric; Glombik, Heiner

PATENT ASSIGNEE(S): Aventis Pharma Deutschland GmbH, Germany

SOURCE: PCT Int. Appl., 36 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 2004087655	A1 20041014	WO 2004-EP2690	20040316
W: AE, AG, AL,	AM, AT, AU, AZ,	BA, BB, BG, BR, BW, BY,	BZ, CA, CH,
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GE, GH, GM,	HR, HU, ID, IL,	IN, IS, JP, KE, KG, KP,	KR, KZ, LC,
LK, LR, LS,	LT, LU, LV, MA,	MD, MG, MK, MN, MW, MX,	MZ, NA, NI,
NO, NZ, OM,	PG, PH, PL, PT,	RO, RU, SC, SD, SE, SG,	SK, SL, SY,
TJ, TM, TN,	TR, TT, TZ, UA,	UG, US, UZ, VC, VN, YU,	ZA, ZM, ZW
RW: BW, GH, GM,	KE, LS, MW, MZ,	SD, SL, SZ, TZ, UG, ZM,	ZW, AM, AZ,
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                                             EP 2004-720854
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                                             NO 2005-5001
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PRIORITY APPLN. INFO.:
                                             DE 2003-10314610
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                                                                    20030811
                                             WO 2004-EP2690
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                                                                    20040316
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OTHER SOURCE(S):

MARPAT 141:314568

GI

$$p-C_6H_4$$
  $p-C_6H_4-OMe$ 

$$p-C_6H_4-CH_2-NH-CO = CH_2 = CO - NH = CH_2 = CH_2$$

AB The invention relates to a novel di-Ph azetidinone (I) and its physiol. compatible salts, to a method for its production, to medicaments containing said

compound and to the use of the latter. Said compound is suitable for use for example as a hypolipidemic agent. Thus, dodecanedioic acid was reacted with thionyl chloride followed by MeOH to give a monomethyl ester, which was then reacted with glucamine and deesterified to give the monoamide intermediate (II). II was reacted with the previously known (2S,3R)-1-(4-aminomethylphenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-(4-methoxyphenyl)azetidin-2-one to give I in 32% yield. In in vitro tests on mice, I had ED50 0.005 mg/mouse for 50% reduction of liver 14C-labeled cholesterol. In solubility tests, compared to a similar reference compound, I

had

better solubility in water, at pH's 1.2, 4.5, 6.8, and 8.0, and in both fasted-

(28  $\mu$ g/mL vs 5) and fed-state simulating intestinal fluids (454  $\mu$ g/mL vs 18) (FaSSIF and FeSSIF).

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE. IN THE RE FORMAT

L26 ANSWER 25 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:1128525 HCAPLUS

DOCUMENT NUMBER: 142:69887

Genetic diagnosis of Alzheimer's disease TITLE:

> susceptibility by detection of gene ABCA2 polymorphism Mace, Sandrine; Ricard, Sylvain; Cousin, Emmanuelle;

INVENTOR(S): Pradier, Laurent; Benavides, Jesus;

Deleuze, Jean Francois

Aventis Pharma S.A., Fr.

PATENT ASSIGNEE(S): Fr. Demande, 41 pp. SOURCE:

CODEN: FRXXBL

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION: DATENT NO

		CENT 1				KINI	)	DATE						NO.			ATE		
		2856				A1	-	2004	1224								0030	520	
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	CA	2529	633			AA		2004	1229		CA 2	004-	2529	633		20	0040	517	
	WO	2004	1135	68		A2		2004	1229	1	WO 2	004-	FR15	09		20	0040	517	
	WO	2004	1135	68		A3			0506										
								AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
			CN.	co,	CR.	CU.	CZ,	DE,	DK,	DM,	DZ,	EC,	ĒE,	EG,	ES,	FI,	GB,	GD,	
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		1639												70					
			-											LU,					
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	CN	1809	•	•	•	•		•		•	•		•	7275			•		1110
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The invention relates to a method of diagnosis and prediction of AB Alzheimer's disease. The method is based on the detection of the presence or absence of the polymorphism in the minority allele rs908832 of gene ABCA2. The presence of the polymorphic allele rs908832 of gene ABCA2 indicates that the subject is in the stage of developing Alzheimer's disease or has an increased risk to develop the disease.

REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 26 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2004:2851 HCAPLUS

DOCUMENT NUMBER:

140:59508

TITLE:

Preparation of diphenylazetidinones substituted by

acidic groups as hypolipidemics.

INVENTOR(S):

Jaehne, Gerhard; Frick, Wendelin; Flohr, Stefanie; Lindenschmidt, Andreas; Glombik, Heiner; Kramer,

Werner; Heuer, Hubert; Schaefer,

Hans-Ludwig

PATENT ASSIGNEE(S):

Aventis Pharma Deutschland G.m.b.H., Germany

SOURCE:

PCT Int. Appl., 48 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PAT	CENT 1	NO.					DATE			APP	LICAT	ION 1	NO.			ATE	
	WO.	2004	0008								wo	2003-	EP58	16			0030	604
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			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN	I, MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
			PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK	C, SL,	ΤJ,	TM,	TN,	TR,	TT,	TZ,
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			BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GÇ	Q, GW,	ML,	MR,	NΕ,	SN,	TD,	TG
	DE	1022	7508			A1		2004	0108		DE	2002-	1022	7508		2	0020	619
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	ΑU	2003	2382	10		A1		2004	0106		AU	2003-	2382	10		2	0030	604
	ΕP	1517	891			A1		2005	0330		ΕP	2003-	7355	35		2	0030	604
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			ΙE,	SI,	LT,	LV,						TR,					SK	
•	BR	2003	0118	96		Α						2003-					0030	
		.1662				Α						2003-						
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	NZ	5373	02			Α		2006	0630			2003-						
		2004				A1		2004				2003-					0030	
		2004				Α		2005				2004-					0041	
		2005				Α		2005	0318			2005-					0050	
PRIOR	TIS	APP	LN.	INFO	.:							2002-						
												2002-					0021	
											WO	2003-	EP58	16		W 2	0030	604

OTHER SOURCE(S):

MARPAT 140:59508

GΙ

Title compds. [I; R1-R6 = H, F, Cl, Br, iodo, CF3, NO2, N3, CN, CO2H, AB CO2alkyl, CONH2, CONHalkyl, CO-30-alkylene-(LAG)n, etc.; n = 1-5; ≥1 C of the alkylenes may be replaced by SOO-2, O, CO, CS, CH:CH, C.tplbond.C, alkylimino, phenylimino, alkylphenylimino, etc.; LAG = (CH2)1-10-SO3H, (CH2)0-10-P(O)(OH)2, (CH2)0-10-OP(O)(OH)2, (CH2)0-10CO2H; with provisos], were prepared Thus, 4-[5-(tert-butyldimethylsilyloxy)-5-(4fluorphenyl) -1-(4-methoxyphenyl) -2-(2-oxo-4-phenyloxazolidin-3carbonyl)pentylamino]benzonitrile (preparation given) in Me tert-Bu ether was treated with N,O-bis(trimethylsilyl)acetamide and Bu4NF in THF and the mixture was stirred 2 h at room temperature to give 4-[3-[3-(tertbutyldimethylsilyloxy) -3-(4-fluorophenyl)propyl]-2-(4-methoxyphenyl)-4oxoazetidin-1-yl]benzonitrile. This was converted to 4-[4-[3-[3-(4fluorophenyl) -3-hydroxypropyl] -2-(4-methoxyphenyl) -4-oxoazetidin-1yl]benzylamino]butane-1-sulfonic acid in several steps. inhibited cholesterol uptake by mouse liver with ED50 = 1.0 mg/mouse orally.

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 27 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2004:2850 HCAPLUS

DOCUMENT NUMBER:

140:77013

TITLE:

Preparation of diphenylazetidinones for the treatment

of hyperlipidemia, arteriosclerosis and

hypercholesterolemia

INVENTOR(S):

Jaehne, Gerhard; Frick, Wendelin; Flohr, Stefanie; Lindenschmidt, Andreas; Glombik, Heiner; Kramer,

Werner; Heuer, Hubert; Schaefer,

Hans-Ludwig

PATENT ASSIGNEE(S):

Aventis Pharma Deutschland G.m.b.H., Germany

SOURCE: PCT Int. Appl., 74 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent German

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT	NO.			KIN	D :	DATE			APPL	ICAT:	ION I	NO.		D	ATE	
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WO 2004	8000	04		A1		2003	1231	•	WO 2	003-1	EP58:	15		2	0030	504
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             FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
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     DE 10227506
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PRIORITY APPLN. INFO.:
                                                                A 20020619
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                                            WO 2003-EP5815
                                                               W 20030604
OTHER SOURCE(S):
                        MARPAT 140:77013
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\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

Title compds. I [R1, R2, R3, R4, R5, R6 = (un)substituted alkylene-(LAG)n; n = 1-5; LAG = sugar; amino sugar; amino acid, etc.] and their pharmaceutically acceptable salts were prepared For example, N-alkylation of 1,4-diazabicyclo[2.2.2]octane with benzyl bromide II, e.g., prepared from 1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxypropyl]-4-(4-hydroxyphenyl)-2-azetidinone and 1,2-bisbromomethylbenzene, afforded diphenylazetidinone III. In rat liver chloresterol absorption assays, 26-examples of compds. I exhibited EC50 values ranging from 0.03-<1.0 (mg/mouse), e.g., the EC50 value of diphenylazetidinone III was 0.3. Compds. I are claimed useful for the treatment of hyperlipidemia, arteriosclerosis and hypercholesterolemia.

REFERENCE COUNT:

3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 28 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2004:2849 HCAPLUS

DOCUMENT NUMBER:

140:77012

TITLE:

Preparation of diphenylazetidinones for the treatment

of hyperlipidemia, arteriosclerosis, and

hypercholesterolemia

INVENTOR(S):

Jaehne, Gerhard; Frick, Wendelin; Flohr, Stefanie; Lindenschmidt, Andreas; Glombik, Heiner; Kramer,

Werner; Heuer, Hubert; Schaefer,

Hans-ludwig

PATENT ASSIGNEE(S):

Aventis Pharma Deutschland GmbH, Germany

SOURCE: PCT Int. Appl., 48 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

## PATENT INFORMATION:

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											WO	200	3 - E	EP58	14		W 2	0030	604
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OTHER SOURCE(S): MARPAT 140:77012

GI

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

Title compds. I [R1, R2, R3, R4, R5, R6 = (un) substituted alkylene-(LAG)n; n = 1-5; LAG = sugar; amino sugar; amino acid, etc.] and their pharmaceutically acceptable salts were prepared For example, condensation of benzonitrile II e.g., prepared from 3-[5-(4-fluorophenyl)-5hydroxypentanoyl]-4-phenyloxazolidin-2-one in 4-steps, and hydroxylamine hydrochloride afforded N-hydroxybenzenecarboximidamide III. In rat liver cholesterol absorption assays, 14-examples of compds. I exhibited EC50 values ranging from 0.03-<1.0 (mg/mouse), e.g., the EC50 value of N-hydroxybenzenecarboximidamide III was 0.1. Compds. I are claimed useful for the treatment of hyperlipidemia, arteriosclerosis, and hypercholesterolemia.

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 29 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

2003:696877 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 139:214472

Preparation of 5-alkoxy-3-phenyl-1,3,4-oxadiazol-2(3H)-TITLE:

ones as inhibitors of pancreatic lipase

Schoenafinger, Karl; Petry, Stefan; Mueller, Guenter; INVENTOR (S):

Bauer, Armin; Heuer, Hubert Otto

PATENT ASSIGNEE(S):

Aventis Pharma Deutschland G.m.b.H., Germany

SOURCE:

PCT Int. Appl., 25 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PAT	CENT 1	NO.			KIN	D	DATE									I	ATE	
	₩O	2003	0725			7.1	_	2002	0004								-	0020	214
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								DK,											
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			PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	, SI	Κ,	SL,	ΤJ,	TM,	TN,	TR,	TT,	TZ,
			UA,	UG,	UΖ,	VC,	VN,	ΥU,	ZA,	ZM,	, ZV	N							
		RW:	GH,	GM,	KΕ,	LS,	MW,	MZ,	SD,	SL,	, S2	Z,	TZ,	ŪĠ,	ZM,	ZW,	AM,	AZ,	BY,
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								GA,											
	DE	1020				A1		2003											
	CA	2477	031					2003											
		2003																	
		1480																	
								ES,											
		10.		-	-			RO,				•			•				
	מם	2003																	
		2005						2004 2005											
		1639				A		2005											
		2003									US	20	03-1	3752	47		2	0030	227
		6900						2005											
		2004				Α		2004	1104										
PRIOR	(TI	APP:	LN.	INFO	. :						DΕ	20	02-3	1020	8987		A 2	0020	228
							•				US	20	02-3	3657	06P		P 2	0020	319
											WO	20	03-1	EP14	84		W 2	0030	214
OTHER	80	HIBCE	191 .			марі	ידעכ	129.	21//	72									

OTHER SOURCE(S):

MARPAT 139:214472

$$R^4$$
 $R^3$ 
 $R^2$ 
 $R^2$ 
 $R^3$ 
 $R^2$ 
 $R^3$ 
 $R^2$ 
 $R^3$ 
 $R^3$ 

AB Title compds. [I; R1 = C7-22 alkyl, (substituted by C4-20 alkoxy, C6-10 aryl, C6-10 aryloxy, or C4-12-alkoxy-C2-4-alkoxy) C2-4 alkyl; C7-C20 alkenyl, 3β- cholestan-3-yl; (substituted) Ph; R2-R5 = H, halo, NO2, (substituted) C1-4 alkyl, C1-9 alkyloxy, CF3, trifluoromethoxy, C6-10-aryl-C1-4-alkyloxy, C6-10 aryloxy, C6-10 aryl, C3-8 cycloalkyl, O-C3-8-cycloalkyl], were prepared for treating obesity. Thus, a mixture of 0.84 g [4-(trifluoromethoxy)phenyl]hydrazine, NMP, and pyridine was

dropwise treated with  $0.43~\mathrm{mL}$  dodecyl chloroformate under ice cooling followed by slow heating at room temperature and stirring for 2 h to give  $0.85~\mathrm{mL}$ 

g

5-dodecyloxy-3-(4-trifluoromethoxyphenyl)-1,3,4-oxadiazol-2(3H)-one. The latter inhibited pancreatic lipase (PL) with IC50 =  $0.03~\mu M$ .

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 30 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

2

ACCESSION NUMBER:

2003:173467 HCAPLUS

DOCUMENT NUMBER:

138:215327

TITLE:

Combined preparations, containing aryl-substituted propanolamine derivatives and other active substances

for the treatment of hyperlipidemia

INVENTOR(S):

Glombik, Heiner; Frick, Wendelin; Schaefer,

Hans-Ludwig; Kramer, Werner

PATENT ASSIGNEE(S):

Aventis Pharma Deutschland GmbH, Germany

PCT Int. Appl., 38 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

German

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

	PAT	ENT I	. O <i>l</i>			KINI	)	DATE		i	APPL	ICAT:	I NOI	. OI		D	ATE	
							-									-		<b>-</b>
	WO	2003	0180	59		A2		2003	0306	1	WO 2	002-I	EP89	07		2	00208	309
	WO	2003	0180	59		<b>A</b> 3		2003	1113									
										BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
												EE,						
												KG,						
												MW,						
												SL,						
								ZA,			,	- ,	•		•	•	•	,
	RW: GH, GM, KE, LS, MW, MZ, S										SZ.	TZ.	UG.	ZM,	ZW,	AM.	AZ,	BY,
		2000										CH,						
												PT,						
						•	•					NE,	-					•
	DE	1014				A1						001-				2	0010	322
		1014				A1						001-					0010	
		2457				AA						002-					0020	309
		1420										002-						
												IT,						
		10.										TR,					,	,
	ВÞ	2002										002-					0020	809
												003-					0020	
	JP 2005505538 T2 200502 CN 1638801 A 200507																0020	
		5312				A						002-				_		
		2004						2004				004-					0040	
PRIOR						11		_004				001-					0010	
PKIOK	. 4 4 1	. ALF		1111	• •							001-			-		0010	
												002-					0020	

OTHER SOURCE(S):

MARPAT 138:215327

GI

AB The invention relates to mixts. of substances, containing propanolamine derivs. of formula (I), in which the functional groups have the indicated meanings, their physiol. acceptable salts and physiol. functional derivs. as well as other active substances for the treatment of metabolic disorders especially hyperlipidemia. The combination can also include antidiabetics, antiarthrytics etc. A typical capsule contains 100 mg of the drugs and 400 mg triglyceride mixture form coco fatty acids; other formulations are emulsions, tablets, dragees, and solns. Hamster that were fed with cholesterol-rich feed received orally the drug combination once daily for 10 days. Feces was analyzed for bile acids, blood lipid levels were measured and cholesterol was determined from liver.

L26 ANSWER 31 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

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ACCESSION NUMBER: 2003:173440 HCAPLUS

DOCUMENT NUMBER: 138:215326

TITLE: Combined preparations, containing 1,4-benzothiepine-

1,1-dioxide derivatives and other active substances

for the treatment of hyperlipidemia

INVENTOR(S): Glombik, Heiner; Frick, Wendelin; Schaefer,

Hans-Ludwig; Kramer, Werner

**Hans-Ludwig**; Kramer, Werner

PATENT ASSIGNEE(S): Aventis Pharma Deutschland GmbH, Germany

SOURCE: PCT Int. Appl., 40 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 2

PAT	rent :	NO.			KIN	<b>D</b> 1	DATE		i	APPL	ICAT	ION 1	NO.		D	ATE	
	2002				7.1		2002	0206							-		
WO	2003	OTOO.	44		AI		2003	0306	,	WO Z	002-	EP89	UB		21	0020	809
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
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		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NO,	NZ,	OM,	PH,
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TN,	TR,	TT,	TZ,
		UA,	UG,	UΖ,	VN,	YU,	ZA,	ZM,	ZW								
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑT,	ΒE,	BG,
		CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙT,	LU,	MC,	NL,
		PT,	SE,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,
		ΝE,	SN,	TD,	TG												
DE	1014	0169			A1	;	2003	0306	3	DE 2	001-	1014	0169		2	0010	822
DE	1014	2456			A1	:	2003	0320	i	DE 2	001-	1014	2456		2	0010	831
CA	2457	976			AA	:	2003	0306	(	CA 2	002-	2457	976		2	0020	809

EP	1425	018			A1	:	2004	0609	E	P	2002-	7962	13		2	0020	809
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	SE,	MC,	PT,
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BR	2002	0120	31		Α	;	2004	0803	В	R	2002-	1203	1		2	0020	809
JP	2005	5018	61		T2	:	2005	0120	J	P	2003-	5225	42		2	0020	809
NZ	5312	93			Α	:	2005	0826	N	Z	2002-	5312	93		2	0020	809
NO	2004	0007	02		Α		2004	0519	N	O	2004 -	702			2	0040	218
PRIORITY	APP	LN.	INFO	. :					D	E	2001-	1014	0169		A 2	0010	822
									D	E	2001-	1014	2456		A 2	0010	831
									W	O	2002-	EP89	08		W 2	0020	809

OTHER SOURCE(S):

MARPAT 138:215326

GT

The invention relates to mixts. of substances, containing 1,4-benzothiepine1,1-dioxide derivs. of formula (I), in which the functional groups have
the indicated meanings, their physiol. acceptable salts and physiol.
functional derivs. as well as other active substances for the treatment of
metabolic disorders especially hyperlipidemia. The combinations can also
include antidiabetics, antiarthrytics etc. A typical capsule contains 100
mg of the drugs and 400 mg triglyceride mixture from coco fatty acids; other
formulations are emulsions, tablets, dragees, and solns. Hamster that
were fed with cholesterol-rich feed received orally the drug
combination once daily for 10 days. Feces was analyzed for bile acids,
blood lipid levels were measured and cholesterol was determined from
liver.

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 32 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:170325 HCAPLUS

DOCUMENT NUMBER: 138:215325

TITLE: Combined preparations, containing aryl-substituted

propanol amine derivatives and other active substances

for the treatment of hyperlipidemia

INVENTOR(S): Glombik, Heiner; Frick, Wendelin; Schaefer,

Hans-Ludwig; Kramer, Werner

PATENT ASSIGNEE(S): Aventis Pharma Deutschland G.m.b.H., Germany

SOURCE: Ger. Offen., 10 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

## PATENT INFORMATION:

PA	TENT	NO.			KIN	D	DATE		7	APP	LICAT	ION I	NO.		D	ATE	
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DE	1014	0170			A1		2003	0306	]	DE	2001-	1014	0170		2	0010	822
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WO	2003	0180	59		A2		2003	0306	1	OW	2002-1	EP89	07		2	0020	809
WO	2003	0180	59		A3		2003	1113									
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											, EE,						
											, KG,						
											, MW,						
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-		UA,	UG,	UZ,	VN,	YU,	ZA,	ZM,	ZW			•	•	•		•	•
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ	, TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,
											, CH,						
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EP	1420		•								2002-'				2	0020	809
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		IE,	SI,	LT,	LV,	FI,	RO.	MK.	CY.	ΑL	, TR,	BG.	cz.	EE.	SK	•	•
BR	2002										2002-					0020	809
JP	2005	5055	38		Т2		2005	0224	,	JP :	2003-	5225	74		2	0020	809
					Α		2005	0713	(	CN	2002-	8163	53		2	0020	809
NZ	CN 1638801 NZ 531292						2005	0826	]	NZ	2002-	5312	92		2	0020	809
US	US 2003158094						2003	0821	1	US	2002-2	2258	02		2	0020	822
ZA	ZA 2004000437										2004-4					0040	121
NO	NO 2004000726						2004	0219	]	NO	2004-	726			2	0040	219
	RIORITY APPLN. INFO.:								]	DE	2001-	1014	0170		A 2	0010	822
									]	DE	2001-	1014	2455		A 2	0010	831
									1	WO :	2002-1	EP89	07	1	W 2	0020	809

OTHER SOURCE(S):

MARPAT 138:215325

G]

AB The invention relates to mixts. of substances, containing propanolamine derivs. of formula (I), in which the functional groups have the indicated meanings, their physiol. acceptable salts and physiol. functional derivs. as well as other active substances for the treatment of metabolic disorders especially hyperlipidemia. The combination can also include antidiabetics, antiarthrytics etc. A typical capsule contains 100 mg of

the drugs and 400 mg triglyceride mixture form coco fatty acids; other formulations are emulsions, tablets, dragees, and solns. The inhibition of sodium-dependent uptake of [3H]-taurocholate (TC) into brush border membrane vesicles was measured.

L26 ANSWER 33 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2003:170324 HCAPLUS

DOCUMENT NUMBER:

138:215324

TITLE:

Combined preparations, containing 1,4-benzothiepine-1,1-dioxide derivatives and other active substances

for the treatment of hyperlipidemia

INVENTOR (S):

Glombik, Heiner; Frick, Wendelin; Schaefer,

Hans-Ludwig; Kramer, Werner

PATENT ASSIGNEE(S):

Aventis Pharma Deutschland G.m.b.H., Germany

SOURCE:

Ger. Offen., 10 pp. CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

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DE	1014	0169			A1		2003	0306		DE	2001-	1014	0169		2	0010	822
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WO	2003	0180	24		A1		2003	0306		WO	2002-	EP89	8 0		2	0020	809
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		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC	, EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE	, KG,	ΚP,	KR,	KZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN	, MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
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							ZA,				, ,						
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	sz	, TZ,	UG,	ZM,	ZW,	AT,	BE,	BG,
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							•	-			, CM,		•		•		-
			SN,			•	•	,			,,	•		~ /		•	•
EP	1425	•	•	•			2004	0609		ΕP	2002-	7962	13		2	0020	809
	R:	AT.	BE.	CH.	DE.	DK.	ES.	FR.	GB.	GR	, IT,	LI.	LU.	NL.	SE	MC,	PT.
		•	•	•	•						, TR,		•	•			
BR	2002										2002-					0020	809
	2005										2003-						
	1655	792	-		A						2002-						
	5312	93			A						2002-						
	2003										2002-					0020	
	2004										2003-					0031	
	2004				A		2004			7. D	2004-	559				0040	126
	2004				A		2004			NO	2004-	702			-	0040	218
PRIORIT								• • • •		DE	2001-	1014	0169		A 2	0010	822
LICKII	- 11L I			• •							2001-					0010	
											2002-					0020	
											2002-		-				
																0	

OTHER SOURCE(S):

MARPAT 138:215324

GI

AB The invention relates to mixts. of substances, containing 1,4-benzothiepine-1,1-dioxide derivs. of formula (I), in which the functional groups have the indicated meanings, their physiol. acceptable salts and physiol. functional derivs. as well as other active substances for the treatment of metabolic disorders especially hyperlipidemia. The combination can also include

antidiabetics, antiarthrytics etc. A typical capsule contains 100 mg of the drugs and 400 mg triglyceride mixture form coco fatty acids; other formulations are emulsions, tablets, dragees, and solns. The inhibition of sodium-dependent uptake of [3H]-taurocholate (TC) into brush border membrane vesicles was measured.

L26 ANSWER 34 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:496163 HCAPLUS

DOCUMENT NUMBER: 140:214414

TITLE: Characterization and identification of the

intestinal cholesterol uptake system

AUTHOR(S): Kramer, W.; Girbig, F.; Corsiero, D.; Burger, K.;

Fahrenholz, F.; Glombik, H.; Heuer, H.

CORPORATE SOURCE: Abt. D.G. Stoffwechsel, Aventis Pharma Deutschland,

Frankfurt, D-65926, Germany

SOURCE: Falk Symposium (2003), 129 (Bile Acids: From Genomics

to Disease and Therapy), 147-160 CODEN: FASYDI; ISSN: 0161-5580 Kluwer Academic Publishers

PUBLISHER: Kluwer Academic Publishers
DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

AB A review presents evidence that both cholesterol and cholesterol absorption inhibitors interact at the level of the brush-border membrane with small intestinal enterocytes. A 145 kDa membrane protein is the mol. target for cholesterol absorption inhibitors catalyzing the first step of intestinal cholesterol absorption, the movement of cholesterol from mixed micelles across the brush-border membrane into the enterocyte. During this step, cholesterol interacts not with the 145 kDa, but with an integral 80 kDa membrane protein. ABC G5 and G8 probably pump out free cholesterol across the brush-border membrane after cholesterol has entered the cell via catalysis by the 145 kDa protein.

REFERENCE COUNT: 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 35 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

DATE

ACCESSION NUMBER:

2002:487559 HCAPLUS

DOCUMENT NUMBER:

137:63115

TITLE:

Preparation of diphenylazetidinone derivatives as

hypolipidemic agents

DATE

INVENTOR (S):

Glombik, Heiner; Kramer, Werner; Flohr, Stefanie;

APPLICATION NO.

Frick, Wendelin; Heuer, Hubert; Jaehne, Gerhard; Lindenschmidt, Andreas; Schaefer,

Hans-Ludwig

PATENT ASSIGNEE(S):

Aventis Pharma Deutschland GmbH, Germany

SOURCE: PCT Int. Appl., 67 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

German

KIND

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.

	PATENT NO.																	
	2002050068															0011	211	
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		GM,	HR,	HU,	ID,	IL,	IN,	ıs,	JP,	KE	, KG,	KP,	KR,	KZ,	LC,	LK,	LR,	
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN	, MW,	MX,	MZ,	NO,	NZ,	OM,	PH,	
											, SL,							
							ZM,											
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ	, TZ,	UG,	ZM,	ZW,	ΑT,	ΒE,	CH,	
		CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE	, IT,	LU,	MC,	NL,	PT,	SE,	TR,	
		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ	, GW,	ML,	MR,	NE,	SN,	TD,	TG	
	1006				A1		2002	0627		DE	2000-	1006	4402		2	0001	221	
DE	1015	4520			A1	2003	1002	DE 2000-10064402 DE 2001-10154520							20011107			
CA	2431	985			AA	2002	0627	CA 2001-2431985							20011211			
AU	U 2002019173						2002	0701		AU	2002-	1917	3		2	0011	211	
EE	2003	0023	7		Α										20011211			
EP	P 1345932										2001-							
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	SE,	MC,	PT,	
							RO,	MK,	CY,	AL	, TR							
BR	2001	0164	82		A 20040203				BR 2001-16482						2	0011		
JP	2004	5162	93		T2 20040603				JP 2002-551564									
NZ	5265	92							NZ 2001-526592									
RU	2275	370			C2 20060427				RU 2003-122219									
US	2002	1282	52		A1				2 US 2001-21028 2							0011	219	
US	6498	156			В2		2002											
	2003		92		Α		2004	0419		ZA	2003-	4092			2	0030		
	2003						2004			ZA 2003-4095						0030		
NO	2003	0027	33		Α		2003	0814		NO 2003-2733 HK 2004-102849					2	0030		
HK PRIORIT	1059	936			A1		2006	0127		HK	2004-	1028	49		2	0040		
PRIORIT	Y APP	LN.	INFO	.:							2000-							
										DE	2001-	1015	4520		A 2	0011	107	
										WO	2001-	EP14	532		W 2	0011	211	
OTHER SO	OURCE	(S):			MAR	PAT	137:	6311	5									
GI																		

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

AB The compds. are suited for use e.g. as hypolipidemic drugs. The invention discloses preparation of diphenylazetidinone derivs. such as I [R1, R2, R3, R4,

R5, R6 = C0-C30-alkylene-L {optionally containing O, CO, CH:CH, C.tplbond.C, N(alkyl), N(alkylphenyl), NH}, H, F, Cl, Br, I, CF3, NO2, CN, CO2H, CO2(alkyl), CONH2, CONH(alkyl), CON(alkyl)2, alkyl, alkenyl, alkynyl, O-alkyl, SO2NH2, SO2NH(alkyl) SO2N(alkyl)2, S-(alkyl), SO(alkyl), (un)substituted S(CH2)nPh, SO(CH2)nPh, SO2(alkyl), SO2(CH2)nPh, NH2, NH(alkyl), N(alkyl)2, NH(acyl), (un)substituted Ph, O(CH2)nPh; n = 0-6; L = II; R7, R9, R10 = Me, Et, Pr, butyl; R8 = H, OH, NH2, NH(alkyl)], and their physiol. acceptable salts, for their use as hypolipidemic agents. Thus, 1,2-diphenylazetidinone derivative III-trifluoroacetate (IV) was prepared via a multistep synthetic sequence starting from 7-[3-(3-butyl-7-dimethylamino-3-ethyl-4-hydroxy-1,1-dioxo-2,3,4,5-tetrahydro-1H-benzo[b]thiepin-5-yl)-phenylcarbamoyl]-heptanoic acid and 4-(4-aminomethylphenyl)-1-(4-fluorophenyl)-3-[3-(4-fluorophenyl)-3-hydroxyphenyl]-azetidin-2-one. Azetidinone IV was tested for its cholesterol lowering ability [ED50 = 0.01 mg/mouse].

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 36 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2002:487551 HCAPLUS

DOCUMENT NUMBER:

137:63114

TITLE:

Preparation of diphenylazetidinone derivatives and

their use as hypolipidemic agents

INVENTOR(S):

Glombik, Heiner; Kramer, Werner; Flohr, Stefanie;

Frick, Wendelin; Heuer, Hubert; Jaehne, Gerhard; Lindenschmidt, Andreas; Schaefer,

Hans-Ludwig

PATENT ASSIGNEE(S):

Aventis Pharma Deutschland GmbH, Germany

SOURCE:

PCT Int. Appl., 41 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT NO.					KIN	CIND DATE			APPLICATION NO.							DATE			
WO 2002050060					A1	1 20020627			1	 WO 2	001-	EP14!	533		2	0011	211		
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	DE 10064402								DE 2000-10064402							0001	221		
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AU	2002	0316	88		A5	:	2002	0701		AU 2	002-								
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ΕP	1345	924			A1		2003	0924		EP 2001-991821					20011211				
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JP	2004	5162	89		T2		2004	0603		JP 2	002-	5515!	56		20011211				
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US 2002128253				A1		2002	0912		US 2001-21044						20011219				

09/05/2006

US 6703386	В2	20040309				
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			WO	2001-EP14533	W	20011211

OTHER SOURCE(S):

MARPAT 137:63114

GT

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

The invention discloses preparation of diphenylazetidinone derivs. such as I [R1, R2, R3, R4, R5, R6 = C0-C30-alkylene-L {optionally containing O, CO, CH:CH, C.tplbond.C, N(alkyl), N(alkylphenyl), NH}, H, F, Cl, Br, I, CF3, NO2, CN, CO2H, CO2(alkyl), CONH2, CONH(alkyl), CON(alkyl)2, alkyl, alkenyl, alkynyl, O-alkyl, SO2NH2, SO2NH(alkyl) SO2N(alkyl)2, S-(alkyl), SO(alkyl), (un) substituted S(CH2) nPh, SO(CH2) nPh, SO2(alkyl), SO2(CH2) nPh, NH2, NH(alkyl), N(alkyl)2, NH(acyl), (un)substituted Ph, O(CH2)nPh; n = 0-6; L = II; Rx, Ry, Rz = H, F, Cl, Br, I, CF3, NO2, CN, CO2H, CO2(alkyl), CONH2, O-alkyl], and their physiol. acceptable salts, for their use as hypolipidemic agents. Thus, 1,2-diphenylazetidinone derivative III trifluoroacetate was prepared from 4-(3-aminomethylphenyl)-1-(4fluorophenyl) -3-[3-(4-fluorophenyl) -3-hydroxypropyl] azetidinone via N-acylation with 11-{2-[3-hydroxy-3-phenyl-2-pyridin-2-yl-1-(pyridin-2ylamino)propyl]-phenylcarbamoyl}-undecanoic acid. Azetidinone III was tested for its cholesterol lowering ability [ED50 = 0.003 mg/mouse]. THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT:

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 37 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2002:487523 HCAPLUS

DOCUMENT NUMBER:

137:63113

TITLE:

Method for producing novel 1,2-diphenylazetidinones,

medicaments containing them, and their use for

treating disorders of lipid metabolism

INVENTOR(S):

Glombik, Heiner; Kramer, Werner; Flohr, Stefanie;

Frick, Wendelin; Heuer, Hubert; Jaehne, Gerhard; Lindenschmidt, Andreas; Schaefer,

Hans-Ludwig

PATENT ASSIGNEE(S):

Aventis Pharma Deutschland GmbH, Germany

PCT Int. Appl., 77 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

German

FAMILY ACC. NUM. COUNT:

PATENT	KIN	D :	DATE		APPLICATION NO.							DATE				
WO 2002	A1 20020627			WO 2001-EP14531							20011211					
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PRIORITY APPLN. INFO.:
                                             DE 2000-10064398
                                                                  A 20011026
W 20011211
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                                             US 2001-21502
                                                                  A3 20011219
                         CASREACT 137:63113; MARPAT 137:63113
OTHER SOURCE(S):
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\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

The invention relates to the compds. I [R1, R2, R3, R4, R5, R6 = AB CO-30-alkylene-LAG {optionally containing O, CO, CH:CH, C.tplbond.C, N(C1-6-alkyl), N(C1-6-alkylphenyl), NH}, H, F, Cl, Br, I, CF3, NO2, CN, CO2H, CO2(C1-6-alkyl), CONH, CONH(C1-6-alkyl), CON(C1-6-alkyl)2, C1-6-alkyl, C1-6-alkenyl, C1-6-alkynyl, O-(C1-6-alkyl), SO2NH2, SO2NH(C1-6-alkyl) SO2N(C1-6-alkyl)2, S-(C1-6-alkyl), SO(C1-6-alkyl), (un) substituted S(CH2) nPh, SO(CH2) nPh, SO2(C1-6-alkyl), SO2(CH2) nPh, NH2, NH(C1-6-alkyl), N(C1-6-alkyl)2, NH(C1-6-acyl), (un)substituted Ph, O(CH2) nPh; LAG = sugar residue, di-, tri-, tetrasaccharide, carbohydrate acid, amino sugar, amino acid, oligopeptide (2 - 9 residues), (trialkylammonium)alkyl, OSO3H] and to their physiol. acceptable salts, suitable, for example, as hypolipidemics. Thus, 1,2-diphenylazetidinone II [R10 = CO(CH2)11NHCO(CHOH)4CH2OH] was prepared from (methoxyphenyl)azetidinone II (R10 = H) via N-acylation with 12-[(2,3,4,5,6-pentahydroxyhexanoyl)amino]dodecanoic acid. Azetidinone II was tested for its cholesterol lowering ability [ED50 = 0.003 mg/mouse]. THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 8 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 38 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2001:669829 HCAPLUS

DOCUMENT NUMBER:

135:356303

TITLE:

GΙ

The caspase-derived C-terminal fragment of  $\beta$  APP induces caspase-independent toxicity and triggers selective increase of A $\beta$ 42 in mammalian cells

Krishnam 10/734,787 05/05/2006

AUTHOR(S): Dumanchin-Njock, Cecile; Alves da Costa, Cristine;

Mercken, Luc; Pradier, Laurent; Checler,

Frederic

CORPORATE SOURCE: Institut de Pharmacologie Moleculaire et Cellulaire,

CNRS, Universite de Nice-Sophia Antipolis, Valbonne,

06560, Fr.

SOURCE: Journal of Neurochemistry (2001), 78(5), 1153-1161

CODEN: JONRA9: ISSN: 0022-3042

PUBLISHER: Blackwell Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

AB During its physiopathol. maturation, the β-amyloid precursor protein undergoes several distinct proteolytic events by activities called

secretases. In Alzheimer's disease, the main histol. hallmark

called senile plaque is clearly linked to the overprodn. of the amyloid peptides A $\beta$ 40 and A $\beta$ 42, two highly aggregable  $\beta$  APP

-derived fragments generated by combined cleavages by  $\beta$ - and  $\gamma$ secretases. Recently, an alternative hydrolytic pathway was
described, involving another category of proteolytic activities called

described, involving another category of proteolytic activities called caspases, responsible for the production of a 31 amino acids  $\beta$  APP C-terminal fragment called C31. C31 was reported to lower the viability of N2a cells but the exact mechanisms mediating C31-toxicity remained to be established. Here the authors show that the transient transfection of pSV2 vector encoding C31 lowers by about 80% TSM1 neuronal cells viability. Arguing against a C31-stimulated apoptotic response, the authors demonstrate by combined enzymic and immunol. approaches that C31 expression did not modulate basal or staurosporine-induced caspase 3-like activity and pro-caspase-3 activation. Furthermore, C31 did not modify Bax and p53 expressions, poly-(ADP-ribose)-polymerase cleavage and cytochrome c translocation into the cytosol. However, the authors established that C31 overexpression triggers selective increase of A $\beta$ 42 but not A $\beta$ 40 production by HEK293 cells expressing wild-type BAPP751. Altogether, the authors' data demonstrate that C31 induces

pathogenic  $\beta$  APP maturation pathway by increasing selectively AB42 species in wild type-  $\beta$  APP-expressing

human cells.

REFERENCE COUNT: 45 THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

a caspase-independent toxicity in TSM1 neurons and potentiates the

L26 ANSWER 39 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1999:795803 HCAPLUS

DOCUMENT NUMBER: 132:35625

TITLE: Amino acid containing benzo[b]thiepine 1,1-dioxide

derivatives as hypolipemic agents

INVENTOR(S): Frick, Wendelin; Enhsen, Alfons; Glombik, Heiner;

Heuer, Hubert

PATENT ASSIGNEE(S): Hoechst Marion Roussel Deutschland G.m.b.H., Germany

SOURCE: PCT Int. Appl., 28 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 9

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 9964410 A1 19991216 WO 1999-EP3701 19990528
W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,

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PRIORITY APPLN. INFO.:
                                          AU 1997-23266
                                                            A3 19970311
                                          WO 1999-EP3701
                                                            W 19990528
                                          US 1999-398315
                                                            A1 19990920
OTHER SOURCE(S):
                       MARPAT 132:35625
GT
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## \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

AB Title compds. such as I (mixture of diastereoisomers) were prepared as hypolipemic agents. Thus, I was prepared in 2 sequences from racemic II and Fmoc-D-lys(Boc)-OH, followed by removal of the Fmoc group with Et2NH. I was ≥20 times more active than 3 analogous comparison substances in tests of fecal separation of 14C-taurocholic acid in rats.

PREFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 40 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1999:795802 HCAPLUS

DOCUMENT NUMBER:

132:22884

TITLE:

Preparation of benzothiepine-1,1-dioxides as

hypolipemics

INVENTOR(S):

Frick, Wendelin; Enhsen, Alfons; Glombik, Heiner;

Heuer, Hubert

PATENT ASSIGNEE(S): Hoechst Marion Roussel Deutschland G.m.b.H., Germany

SOURCE: PCT Int. Appl., 30 pp.

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE: Patent German

FAMILY ACC. NUM. COUNT: 9

PAT	ENT NO.	KIND	DATE	APPLICATION NO.	DATE			
WO	9964409 9964409	A2	19991216	WO 1999-EP3743	19990529			
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CN	1127497	В	20031112	CN 1999-807171	19990528			
CA	2334773	AA	19991216	CA 1999-2334773	19990529			
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US 1999-398315 A1 19990920 US 2001-773772 A1 20010202 US 2002-201050 A1 20020724

OTHER SOURCE(S):

MARPAT 132:22884

GΙ

$$\mathbb{R}^{4}\mathbb{R}^{5}\mathbb{N}$$

AB Title compds. [I; R = C6H4NHZR3; R1,R4,R5 = Me, Et, Pr, Bu; R2 = H, OH, amino(alkyl); R3 = sugar residue; Z = bond, carbonyl(alkylene), CONH, etc.] were prepared Thus, I [R = C6H4(NHR')-3, R1 = Et, R2 = OH, R4 = R5 = Me](II; R' = H) was amidated by penta-O-acetyl-D-gluconic acid and the product deprotected to give II (R' = gluconoyl) as a mixture of diastereomers. Data for biol. activity of I were given.

L26 ANSWER 41 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

Ι

ACCESSION NUMBER: 1999:602051 HCAPLUS

DOCUMENT NUMBER: 131:347939

TITLE: Substrate specificity of the ileal and the hepatic

Na+/bile acid cotransporters of the rabbit. I.
Transport studies with membrane vesicles and cell

lines expressing the cloned transporters

AUTHOR(S): Kramer, Werner; Stengelin, Siegfried; Baringhaus,

Karl-Heinz; Enhsen, Alfons; Heuer, Hubert;

Becker, Wolfgang; Corsiero, Daniel; Girbig, Frank;

Noll, Rudiger; Weyland, Claudia

CORPORATE SOURCE: DG Metabolic Diseases, Hoechst Marion Roussel

Deutschland GmbH, Frankfurt am Main, D-65926, Germany

SOURCE: Journal of Lipid Research (1999), 40(9), 1604-1617

CODEN: JLPRAW; ISSN: 0022-2275

PUBLISHER: Lipid Research, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

ΔR The substrate specificity of the ileal and the hepatic Na+/bile acid cotransporters was determined using brush border membrane vesicles and CHO cell lines permanently expressing the Na+/bile acid cotransporters from rabbit ileum or rabbit liver. The hepatic transporter showed a remarkably broad specificity for interaction with cholephilic compds. in contrast to the ileal system. The anion transport inhibitor diisothiocyanostilbene disulfonate (DIDS) is a strong inhibitor of the hepatic Na+/bile acid cotransporter, but does not show any affinity to its ileal counterpart. Inhibition studies and uptake measurements with about 40 different bile acid analogs differing in the number, position, and stereochem. of the hydroxyl groups at the steroid nucleus resulted in clear structure-activity relationships for the ileal and hepatic bile acid transporters. The affinity to the ileal and hepatic Na+/bile acid cotransport systems and the uptake rates by cell lines expressing those transporters as well as rabbit ileal brush border membrane vesicles is primarily determined by the substituents on the steroid

nucleus. Two hydroxy groups at position 3, 7, or 12 are optimal whereas the presence of three hydroxy groups decreased affinity. Vicinal hydroxy groups at positions 6 and 7 or a shift of the 7-hydroxy group to the 6-position significantly decreased the affinity to the ileal transporter in contrast to the hepatic system. 6-Hydroxylated bile acid derivs. are preferred substrates of the hepatic Na+/bile acid cotransporter. Surprisingly, the 3\alpha-hydroxy group being present in all natural bile acids is not essential for high affinity interaction with the ileal and the hepatic bile acid transporter. The  $3\alpha$ -hydroxy group seems to be necessary for optimal transport of a bile acid across the hepatocyte canalicular membrane. A modification of bile acids at the 3-position therefore conserves the bile acid character thus determining the 3-position of bile acids as the ideal position for drug targeting strategies using bile acid transport pathways.

REFERENCE COUNT:

55

THERE ARE 55 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 42 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1997:812258 HCAPLUS

DOCUMENT NUMBER:

128:57465

TITLE:

Use of inhibitors of cellular

Na+/H+-exchangers for preparing a medicine for

normalizing serum lipids

INVENTOR (S):

Lang, Hans Jochen; Schwark, Jan Robert; Kleemann,

Heinz Werner; Jung, Oliver; Schaefer, Hans

Ludwig; Linz, Wolfgang; Kramer, Werner;

Schoelkens, Bernward; Jansen, Hans Willi; Falk, Eugen

PATENT ASSIGNEE(S):

Hoechst A.-G., Germany

SOURCE: Ger. Offen., 6 pp. CODEN: GWXXBX

Patent

DOCUMENT TYPE: LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT NO.						1	APPL	ICAT:	ION I		DATE								
DE	1962		A1			:													
CA	2257	299			AA		1997	1211		CA 1	997-2	22572	299		1:	9970!	520		
WO	9746							1211	Ĭ	WO 1	997-1	EP254	48		1	9970	520		
WO	9746				A3														
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		IL,	IS,	JP,	KG,	KΡ,	KR,	ΚZ,	LC,	LK,	LR,	LT,	LV,	MD,	MG,	MK,	MN,		
		MX,	NO,	ΝZ,	PL,	RO,	RU,	SG,	SI,	SK,	ТJ,	TM,	TR,	TT,	UA,	US,	UΖ,		
		VN,																	
	RW:	GH,																	
		GR,	ΙE,	IT,	LU,	MC,	ΝL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,		
					SN,														
AU 9729576													19970520						
	7221																		
									EP 1997-923937						19970520				
EΡ	9185																		
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CN	1221	339			Α		1999	0630	1	CN 1	997-	1951	94						
BR	9709	516			Α	:	1999	0810								9970!	520		
	2000							0606			998-				19970520				
NZ	3330	95			Α	:	2000	0825	1	NZ 1	997-3	3330	95		1:	19970520			
	2211				C2										19970520				
AT	2939	65			Е	:	2005	0515		AT 1	997-	9239:	37	19970520					

AB Na+/H+ exchangers (especially guanidine derivs.) are useful in medications for lowering serum lipid levels, treatment of hypercholesterolemia-related circulatory disorders, and prevention and treatment of atherosclerosis, endothelial dysfunction syndrome, cardiac hypertrophy, cardiomyopathy, coronary vasospasm, and myocardial infarct and of disorders secondary to these diseases. Thus, in rabbits receiving a cholesterol-rich diet, addition of 0.1% HOE 642 [(4-isopropyl-3-methanesulfonyl)benzoylguanidi ne methanesulfonate] to the diet decreased the serum cholesterol, VLDL, LDL, and HDL levels from 17.85, 5.68, 9.31, and 2.86 (control) to 7.95, 4.6, 2.1, and 1.8 mM, resp.

L26 ANSWER 43 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1997:161940 HCAPLUS

TITLE:

Synthesis and SAR of trimeric bile acid reabsorption

inhibitors: A new approach to lower

cholesterol

AUTHOR (S):

Glombik, H.; Baringhaus, K. -H.; Boeger, G.; Enhsen, A.; Falk, E.; Friedrich, M.; Hoffmann, A.; Kramer, W.;

Schaefer, H. L.; et al.

CORPORATE SOURCE:

HMR TA Metabolism Research, Frankfurt/Main, D-65926,

Germany

SOURCE:

Book of Abstracts, 213th ACS National Meeting, San Francisco, April 13-17 (1997), MEDI-108. American

Chemical Society: Washington, D. C.

CODEN: 64AOAA

DOCUMENT TYPE:

Conference; Meeting Abstract

LANGUAGE: English

AB Recent attempts in antiatherosclerotic therapy focus on cholesterol lowering agents with new modes of action. With regard to resins and statins real progress is expected from non absorbable bile acid reabsorption inhibitors (BARI) that block the ileal transporter specific for bile acids. This will lead to increased excretion of bile acids, resynthesis from cholesterol in the liver and thus lowering of blood cholesterol by an indirect and non systemic mechanism. While there is some information available on the size and function of the ileal bile acid transporter, a detailed structure anal. of this transmembrane protein has not been performed. BARI with minimal absorption are designed and synthesized by combining bile acid moieties as recognition units via linkers to trivalent core structures such as Kemp's triacid. Linker chemical had to be developed for this purpose. A directing effect of the core unit is most important for activity at the primary target as tested in cell and animal models.

L26 ANSWER 44 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1995:308716 HCAPLUS

DOCUMENT NUMBER:

122:81416

TITLE:

Heterocycle-containing amidine derivatives, their

preparation, and use as LTB4 antagonists

INVENTOR(S): Renth, Ernst Otto; Schromm, Kurt; Anderskewitz, Ralf;

Birke, Franz; Fuegner, Armin; Heuer, Hubert

PATENT ASSIGNEE(S): Boehringer Ingelheim International G.m.b.H., Germany

SOURCE: Ger. Offen., 31 pp.

CODEN: GWXXBX

DOCUMENT TYPE:

LANGUAGE:

GI

Patent German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT																			
						-													
DE	DE 4309285				A1		19940929			DE 1993-4309285					19930323				
CA	CA 2158994				AA		19940929			CA 1994-2158994					19940318				
								WO 1994-EP856											
	W:	AT,	AU,	ВG,	BY,	CA,	CH,	CN,	CZ,	DE,	DK,	ES,	FI,	GB,	HU,	JP,	KR,		
		ΚZ,	LU,	LV,	NL,	NO,	NZ,	PL,	PT,	RO,	RU,	SE,	SK,	UA,	US				
	RW:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE		
AU	9463780				A1	A1 19941011				AU 1994-63780					19940318				
EP	690849				A1		1996	EP 1994-911191				19940318							
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,	LI,	LU,	MC,	NL,	PT,	SE	
CN	1124486				Α		1996	0612	(	CN 1994-192207				19940318					
JP	JP 08508467				T2	T2 19960910				JP 1994-520657				19940318					
HU 73968				A2	A2 19961028				HU 1995-2778					19940318					
ZA	ZA 9401993				Α	A 19940923				ZA 1994-1993					19940322				
FI	FI 9504491				A 19950922				FI 1995-4491										
NO									NO 1995-3763										
LV	1146							1220								9950			
PRIORITY		_			_				_				285						
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OTHER SOURCE(S):					MARI	PAT	122:	8141				21.00	•						

$$\begin{array}{c|c} R^4 \\ \hline \\ Het - A \\ \hline \\ B \\ \hline \\ NH_2 \\ I \\ \end{array}$$

II

Searched by Paul Schulwitz 571-272-2527

treating inflammatory and allergic conditions such as asthma, ulcerative

colitis, psoriasis, and gastropathy induced by nonsteroidal antiphlogistics. For example, Pinner reaction of 4-[2-(2-benzothiazolyloxy)ethoxy]benzonitrile, by treatment with HCl and EtOH in CH2Cl2 at -15°, and ammonolysis of the precipitated crystalline imidate with NH3-saturated EtOH at reflux, gave title compound II as the HCl salt. I had Ki of 1-20 nM in an LTB4 receptor binding assay. Over 90 I are listed with m.p. data.

L26 ANSWER 45 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1987:211507 HCAPLUS

DOCUMENT NUMBER: 106:211507

TITLE: Ion transport and electrophysiology of the early

proximal colon of rabbit

AUTHOR(S): Clauss, W.; Biehler, K. H.; Schaefer, H.;

Wills, N. K.

CORPORATE SOURCE: Inst. Zoophysiol., Univ. Hohenheim, Stuttgart, D-7000,

Fed. Rep. Ger.

SOURCE: Pfluegers Archiv (1987), 408(6), 592-9

CODEN: PFLABK; ISSN: 0031-6768

DOCUMENT TYPE: Journal LANGUAGE: English

Transepithelial transport of Na+, K+, and Cl- in the isolated initial segment (P1) of rabbit colon in vitro was studied by using radioisotopic tracer fluxes and electrophysiol. techniques. Like the rabbit descending colon, the proximal colon actively absorbs Na+ and Cl-; however, its transport systems are markedly different. In vivo, this segment absorbs K+, but active K+ secretion was observed in vitro. Unlike the descending colon, Na+ absorption is relatively insensitive to amiloride, and only a slight inhibition was obtained even at 1 mM concns. of this drug. Na+ and Cl- absorption appeared to be coupled (directly or indirectly), since the absorption of each ion was inhibited by the removal of the other. Serosal ouabain also inhibited Na+ and Cl- absorption and net K+ secretion. Unlike the descending colon, the proximal P1 segment did not have a net absorptive K+ transport system that was detectable in the presence of ouabain. Elec., the early proximal colon has a low transepithelial resistance compared to descending colon (RT = 133  $\Omega/\text{cm2}$ ) but a larger short-circuit current (Isc = 178  $\mu A/cm2$ ). The transepithelial potential averaged -21 mV, in excellent agreement with values measured in vivo. The apical and basolateral membrane potentials averaged -21 mV and -42 mV, and intracellular K+ activity was 70 mM. The findings indicate active K+ uptake across the basolateral membrane and passive exit across the apical membrane. The basolateral membrane conductance may be a K+ conductance that is blockable by Ba+. It is likely that K+ transport normally occurs by both cellular and paracellular routes in this epithelium. Because of the numerous differences between this segment and the descending colon, the P1 segment of proximal colon apparently has a distinct function in colonic electrolyte transport.

L26 ANSWER 46 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1985:39549 HCAPLUS

DOCUMENT NUMBER: 102:39549

TITLE: Cationophore properties of the new polyether

antibiotic salinomycin investigated in distal rabbit

colon in vivo and in vitro

AUTHOR(S): Schaefer, H.; Clauss, W.; Hoernicke, H.

CORPORATE SOURCE: Inst. Zoophysiol., Univ. Stuttgart-Hohenheim,

Stuttgart, D-7000/70, Fed. Rep. Ger.

SOURCE: Comparative Biochemistry and Physiology, Part A:

Molecular & Integrative Physiology (1984), 79A(3),

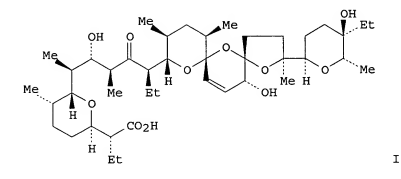
387-92

CODEN: CBPAB5; ISSN: 0300-9629

DOCUMENT TYPE: LANGUAGE: Journal

GΙ

English



AB In the distal rabbit colon, salinomycin (I) [53003-10-4] (104 M) did not influence net water, Cl- or, Na+ absorption, but it decreased K+ secretion to zero and the transepithelial potential was decreased from -45 mV to -33 mV. I at 10-4 and 10-3 M applied to the mucosal side of the colon decreased the transepithelial potential from 18 mV to zero within 80 and 30 min, resp. I also affected the short-circuit current and the transepithelial conductance in a dose-dependent manner. The unidirectional 22Na fluxes were increased to 20 times the control values and the net Na transport was inhibited by I. Thus, I given in doses used as a coccidiostatic feed additive profoundly affected colon electrolyte transport.

L26 ANSWER 47 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1985:106108 HCAPLUS

DOCUMENT NUMBER:

102:106108

TITLE:

Indalpine, a potent and selective 5-HT uptake blocker

AUTHOR (S):

Le Fur, G.; Gueremy, C.; Benavides, J.;

Malgouris, C.; Uzan, A.

CORPORATE SOURCE:

Pharm. Lab., Groupe Rhone Poulenc Sante,

Gennevilliers, Fr.

SOURCE:

Advances in Biological Psychiatry (1984), 14 (Serotonin

Affective Disord.), 33-40, 1 plate

CODEN: ABPSD5; ISSN: 0378-7354

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB 5-HT [50-67-9] uptake by rat brain in vitro was inhibited by indalpine (I) [63758-79-2]; I showed no affinity for catecholamine or amino acid uptake systems. Twice daily administration of I (10 mg/kg, i.p.) during a 14 day period did not downregulate β, 5-HT1, or 5-HT2 receptors. Thus the therapeutic effect of I is not associated with downregulation of monoamine postsynaptic receptors. In binding studies, the inhibition of I binding to rat brain by tricyclic antidepressants paralleled their inhibition of 5-HT uptake. No correlation, however, was found between the inhibition of I binding and inhibition of noradrenaline uptake by antidepressants. I binding in rat brain was competitively inhibited by 5-HT. Pharmacol. studies of I binding sites in brain

suggest these sites may be located in 5-HT neurons. Thus, I is a potent 5-HT reuptake blocker characterized by a very high selectivity of action due to its affinity to presynaptic sites.

L26 ANSWER 48 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1975:69569 HCAPLUS

DOCUMENT NUMBER: 82:69569

TITLE: Estimation of enzyme activity in living epidermal

cells

AUTHOR(S): Schalla, W.; Zesch, A.; Schaefer, H.

CORPORATE SOURCE: Rudolf-Virchow-Hosp., Free Univ., Berlin, Fed. Rep.

Ger.

SOURCE: British Journal of Dermatology (1974), 91(5), 489-501

CODEN: BJDEAZ; ISSN: 0007-0963

DOCUMENT TYPE: Journal LANGUAGE: English

Amethod is described for measurement of cytoplasmic enzymes in intact cells of excised human epidermis. The method is suitable for determination of lactic dehydrogenase (I), malate dehydrogenase (II), pyruvate kinase (III), glutamate-oxalacetate transaminase, and NADP-dependent isocitrate dehydrogenase, but not of mitochondrial glutamate dehydrogenase and NAD-dependent isocitrate dehydrogenase. Ouabain inhibited activity of cellular but not of extracellular I, II, and III. Fluoromalate inhibited activity of both forms of II. The activity of I was higher under anaerobic than under aerobic conditions. The app. can be modified for determining enzyme activity in epidermis in situ, following removal of stratum corneum by tape stripping. Activity of I, but not of II, is significantly higher in psoriatic than in normal skin. Activity of both enzymes is reduced within 15 min by oral

L26 ANSWER 49 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1970:130964 HCAPLUS

DOCUMENT NUMBER: 72:130964

TITLE: Biochemical studies of psoriasis chemotherapy

AUTHOR(S): Schaefer, H. CORPORATE SOURCE: Fed. Rep. Ger.

SOURCE: Archiv fuer Klinische und Experimentelle Dermatologie

administration of prednisolone or i.v. administration of methotrexate.

(1968), 237, 240-5

CODEN: AKEDAX; ISSN: 0300-8614

DOCUMENT TYPE: Journal LANGUAGE: German

AB The inhibitory effect of Cignolin (I) and 6-hydroxy-1,3-

benzoxathiol-2-one (II) on cutaneous enzymes was investigated. I inhibited glucose-6-phosphate dehydrogenase and reacted with

NADPH. II inhibited glucose-6-phosphate dehydrogenase,

glutamic-oxalacetic transaminase, malic dehydrogenase and leucine aminopeptidase, but not aldolase, lactic dehydrogenase, isocitric dehydrogenase, and glutathione reductase.

L26 ANSWER 50 OF 65 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1949:50774 HCAPLUS

DOCUMENT NUMBER: 43:50774
ORIGINAL REFERENCE NO.: 43:9108b-f

TITLE: Critique and procedure for cholinesterase

determinations in blood

AUTHOR(S): Schaefer, Hans; Maier, Erich

SOURCE: Biochemische Zeitschrift (1949), 319, 420-38

CODEN: BIZEA2; ISSN: 0366-0753

DOCUMENT TYPE:

LANGUAGE:

Unavailable

The hydrolysis of acetylcholine (ACh) in human blood under physiol. conditions, i.e., very small concns., is almost entirely due to cholinesterase of the erythrocytes and practically none to cholinesterase of the serum. Therefore, changes in serum cholinesterase values are physiologically without importance so long as the erythrocyte cholinesterase values are normal. It is important to bear clearly in mind that, wherever cholinesterase is diminished, the ACh liberated at any parasympathetic ending remains active for a longer than normal duration. As a result a vagotonus develops, or temporary predominant parasympathetic innervation. Of course, this hypothesis presupposes that the ACh measured is the ACh which is effective in the vegetative field upon which the parasympathetic acts. It always seemed more or less doubtful whether the serum cholinesterase represented such a reference. The motor end plate is definitely the place where ACh is liberated, as can be judged by its high local concentration. It is not permissible thus to regard a decrease in serum cholinesterase as an indication of increased vagotonus. Besides, since the serum cholinesterase is presumably an unspecific pseudocholinesterase, its variations probably reflect changes in the composition of protein fractions rather than those in the vegetative hormonal system. But neither does the determination of erythrocyte cholinesterase reveal anything regarding the cholinergic transmission at the vegetative end organs. Certain kinetic constants must be measured to determine the cholinesterase activity of erythrocytes. This is done in the Warburg app. and from these detns. the relative cholinesterase concentration is calculated, as well as the mode

of binding of ACh and cholinesterase, and finally the equilibrium constant of the inhibitory reaction between cholinesterase and ACh. The cholinesterase concentration attains a min. at about 30 years of age.

L26 ANSWER 51 OF 65 ACCESSION NUMBER:

MEDLINE on STN

DOCUMENT NUMBER:

2005184964 MEDLINE PubMed ID: 15816856

TITLE:

Expression of human FE65 in amyloid precursor protein

transgenic mice is associated with a reduction in

beta-amyloid load.

AUTHOR:

Santiard-Baron Dominique; Langui Dominique; Delehedde Maryse; Delatour Benoit; Schombert Brigitte; Touchet Nathalie; Tremp Gunter; Paul Marie-Francoise; Blanchard Veronique; Sergeant Nicolas; Delacourte Andre; Duyckaerts

Charles: Pradier Laurent: Mercken Luc

CORPORATE SOURCE:

Neurodegenerative Diseases Group, Aventis, Vitry-sur-Seine,

France.

SOURCE:

Journal of neurochemistry, (2005 Apr) Vol. 93, No. 2, pp.

Journal code: 2985190R. ISSN: 0022-3042.

PUB. COUNTRY:

England: United Kingdom

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

200505

ENTRY DATE:

Entered STN: 9 Apr 2005

Last Updated on STN: 21 May 2005 Entered Medline: 20 May 2005

AB FE65 is an adaptor protein that interacts with the cytoplasmic tail of the amyloid precursor protein (APP). In cultured non-neuronal cells, the formation of the FE65-APP complex is a key element for the modulation of APP processing, signalling and

beta-amyloid (Abeta) production. The functions of FE65 in vivo, including its role in the metabolism of neuronal APP, remain to be investigated. In this study, transgenic mice expressing human FE65 were generated and crossbred with APP transgenic mice, known to develop Abeta deposits at 6 months of age. Compared with APP mice, APP/FE65 double transgenic mice exhibited a lower Abeta accumulation in the cerebral cortex as demonstrated by immunohistochemistry and immunoassay, and a lower level of APP The reduced accumulation of Abeta in APP/FE65 double transgenics, compared with APP mice, could be linked to the low Abeta42 level observed at 4 months of age and to the lower APP -CTFs levels. The present work provides evidence that FE65 plays a role in the regulation of APP processing in an in vivo model.

L26 ANSWER 52 OF 65 MEDLINE on STN ACCESSION NUMBER: 91264881 MEDLINE DOCUMENT NUMBER: PubMed ID: 1646613

TITLE: Pharmacodynamics, pharmacokinetics and safety profile of the new platelet-activating factor antagonist apafant in

AUTHOR: Brecht H M; Adamus W S; Heuer H O; Birke F W;

Kempe E R

CORPORATE SOURCE: Department of Medicine, Boehringer Ingelheim KG, Rhein,

Germany.

SOURCE: Arzneimittel-Forschung, (1991 Jan) Vol. 41, No. 1, pp.

51-9.

Journal code: 0372660. ISSN: 0004-4172.

PUB. COUNTRY: GERMANY: Germany, Federal Republic of

DOCUMENT TYPE: (CLINICAL TRIAL)

Journal; Article; (JOURNAL ARTICLE)

(RANDOMIZED CONTROLLED TRIAL)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199107

ENTRY DATE: Entered STN: 2 Aug 1991

> Last Updated on STN: 2 Aug 1991 Entered Medline: 15 Jul 1991

Platelet-activating factor (PAF) is a unique phospholipid mediator with AB multifunctional properties. Evidence generated in experimental studies suggests that PAF plays a pathogenetic role in anaphylactic, inflammatory and immunogenic reactions. Apafant (WEB 2086, CAS 105219-56-5), a novel synthetic PAF receptor antagonist, was administered to a total of 101 healthy volunteers within 5 studies to investigate its pharmacologic activity, pharmacokinetic behaviour and safety profile. Pharmacologic activity was monitored by inhibition of 5 x 10(-8) mol/1 PAF-induced platelet aggregation ex vivo. The following treatment schedules were studied: oral single dose 1.25 to 400 mg; oral multiple dose 100 mg t.i.d. over 7 days; i.v. infusion 0.5 to 50 mg (over 30 min); inhalative administration up to 1.0 mg. PAF induced platelet aggregation was virtually completely inhibited by single oral doses of 20 mg upwards, throughout during the multiple oral dose study, at all dose levels tested in the i.v. study and (significantly but not completely) at 0.5 and 1.0 mg in the inhalative study. Following oral administrations (capsules) apafant is absorbed rapidly (tmax 1 to 2 h), there is linear pharmacokinetics for the mean plasma concentrations of apafant measured by RIA as well as for the areas under the curve (AUCs). Approximately 60% of apafant is bound to plasma protein, the mean volume of distribution is 28 1, about 44% of an oral dose is excreted in the urine, the mean renal clearance is 192 ml/min. No accumulation of the drug occurred in

volunteers with normal kidney function. No clinically relevant drug related adverse events or changes in laboratory or vital parameters such as blood pressure, heart rate, respiratory rate and ECG were observed. (ABSTRACT TRUNCATED AT 250 WORDS)

L26 ANSWER 53 OF 65 EMBASE COPYRIGHT (c) 2006 Elsevier B.V. All rights

reserved on STN

90126142 EMBASE ACCESSION NUMBER:

DOCUMENT NUMBER:

1990126142

TITLE:

Tumor necrosis factor (TNF) and endotoxin prime effects of

PAF in vivo.

AUTHOR:

Heuer H.O.; Letts G.; Meade C.J.

CORPORATE SOURCE:

Department of Pharmacology, Boehringer Ingelheim, D-6507

Ingelheim/Rhein, Germany

SOURCE:

Journal of Lipid Mediators, (1990) Vol. 2, No. SUPPL., pp.

S101-S108.

ISSN: 0921-8319 CODEN: JLMEEG

COUNTRY:

Netherlands

DOCUMENT TYPE: FILE SEGMENT:

Journal; Conference Article

025 Hematology

Immunology, Serology and Transplantation 026

Drug Literature Index 037

LANGUAGE:

English English

SUMMARY LANGUAGE: ENTRY DATE:

Entered STN: 13 Dec 1991

Last Updated on STN: 13 Dec 1991

The purpose of the present study in NMRI mice was to investigate the AB action of platelet-activating factor (PAF) on mortality and intestinal transit velocity, the interaction of endotoxin or tumor necrosis factor (TNF) with the effect of PAF on these parameters and the effect of the PAF antagonist WEB 2086 on the endotoxin/TNF- and PAF-induced changes. PAF at a high dose (200 μg/kg i.v.) increased mortality and reduced transit velocity. This effect was inhibited by WEB 2086 (0.01 - 0.5 mg/kg i.p.) in a dose-dependent manner. Pretreatment with endotoxin (S. typhosa; 10 µg/kg i.v.) or TNF (40 μq/kg i.v.) enhanced the activity of PAF resulting in increased mortality and reduced transit velocity. This enhanced activity of PAF in the case of pretreatment with endotoxin or TNF occurred at doses at which PAF, endotoxin or TNF given alone did not significantly affect these parameters. The ability of endotoxin or TNF to enhance the effect of PAF was maximal, if the time delay between endotoxin and subsequent PAF administration was about 1 - 2 h. WEB 2086 (0.01 - 1 mg/kg i.p.) inhibited this priming in a dose-dependent fashion. These findings support suggestions of a role for PAF in endotoxin shock and TNF-associated shock-like syndrome.

L26 ANSWER 54 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

STN

ACCESSION NUMBER: 2004:210443 BIOSIS DOCUMENT NUMBER: PREV200400213329

TITLE:

Diphenylazetidinone derivatives, process for their preparation, medicaments comprising these compounds and

AUTHOR (S):

Glombik, Heiner [Inventor, Reprint Author]; Kramer, Werner [Inventor]; Flohr, Stefanie [Inventor]; Frick, Wendelin

[Inventor]; Heuer, Hubert [Inventor]; Jaehne,

Gerhard [Inventor]; Lindenschmidt, Andreas [Inventor];

Schaefer, Hans-Ludwig [Inventor]

CORPORATE SOURCE:

Hofheim, Germany

ASSIGNEE: Aventis Pharma Deutschland GmbH, Frankfurt am

Main, Germany

PATENT INFORMATION: US 6703386 20040309

Official Gazette of the United States Patent and Trademark SOURCE:

Office Patents, (Mar 9 2004) Vol. 1280, No. 2. http://www.uspto.gov/web/menu/patdata.html. e-file.

ISSN: 0098-1133 (ISSN print).

DOCUMENT TYPE:

Patent English

LANGUAGE: ENTRY DATE:

Entered STN: 14 Apr 2004

Last Updated on STN: 14 Apr 2004

Compounds of the formula I, for example, are disclosed, ##STR1## in which AB R1, R2, R3, R4, R5, and R6 independently of one another are (C0

-C30)-alkylene-L or are the meanings given in the description, and where L is shown connected to (C0 -C30)-alkylene as follows: ##STR2## where Rx,

Ry, Rz have the meanings given in the description, and their

physiologically acceptable salts. The compounds are suitable for use, for example, as hypolipidemics.

L26 ANSWER 55 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

STN

ACCESSION NUMBER:

2006:158029 BIOSIS PREV200600174128

DOCUMENT NUMBER: TITLE:

The squalene synthase inhibitor RPR 107393A

reduces A beta peptide level in an APP transgenic

mouse model.

AUTHOR (S):

Pradier, Laurent [Reprint Author]; Canton, Thierry; Bouaiche, Zakia; Benoit, Patrick;

Benavides, Jesus

CORPORATE SOURCE:

Aventis Pharma, Vitry Sur Seine, France

laurent.pradier@aventis.com

SOURCE:

Neurobiology of Aging, (JUL 2004) Vol. 25, No. Suppl. 2,

pp. S568.

Meeting Info.: 9th International Conference on Alzheimers Disease and Related Disorders. Philadelphia, PA, USA. July

17 -22, 2004. Alzheimers Assoc. CODEN: NEAGDO. ISSN: 0197-4580.

DOCUMENT TYPE:

Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LANGUAGE:

English

ENTRY DATE:

Entered STN: 9 Mar 2006

Last Updated on STN: 9 Mar 2006

L26 ANSWER 56 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

STN

ACCESSION NUMBER:

2006:156619 BIOSIS

DOCUMENT NUMBER:

PREV200600172718

TITLE:

The role of the tissue-type plasminogen activator in the

course of amyloid accumulation.

AUTHOR (S):

Cacquevel, Mathias [Reprint Author]; Cheenne, Simon;

Castel, Herve; Benavides, Jesus; Pradier,

Laurent; Vivien, Denis

CORPORATE SOURCE:

Aventis Pharma, Vitry Sur Seine, France

m.cacquevel@cyceron.fr

SOURCE:

Neurobiology of Aging, (JUL 2004) Vol. 25, No. Suppl. 2,

pp. S141.

Meeting Info.: 9th International Conference on Alzheimers Disease and Related Disorders. Philadelphia, PA, USA. July

17 -22, 2004. Alzheimers Assoc.

Krishnan 10/734,787

CODEN: NEAGDO. ISSN: 0197-4580.

DOCUMENT TYPE: Conference; (Meeting)

Conference; (Meeting Poster)

LANGUAGE: English

ENTRY DATE: Entered STN: 9 Mar 2006

Last Updated on STN: 9 Mar 2006

L26 ANSWER 57 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

STN

ACCESSION NUMBER: 2003:83632 BIOSIS DOCUMENT NUMBER: PREV200300083632

TITLE: Diphenylazetidinone derivatives, process for their

preparation, medicaments comprising these compounds and

. .

their use.

AUTHOR(S): Glombik, Heiner [Inventor, Reprint Author]; Kramer, Werner

[Inventor]; Flohr, Stefanie [Inventor]; Frick, Wendelin

[Inventor]; Heuer, Hubert [Inventor]; Jaehne,

Gerhard [Inventor]; Lindenschmidt, Andreas [Inventor];

Schaefer, Hans-Ludwig [Inventor]

CORPORATE SOURCE: Hofheim, Germany

ASSIGNEE: Aventis Pharma Deutschland GmbH, Frankfurt am

Main, Germany

PATENT INFORMATION: US 6498156 20021224

SOURCE: Official Gazette of the United States Patent and Trademark

Office Patents, (Dec 24 2002) Vol. 1265, No. 4. http://www.uspto.gov/web/menu/patdata.html. e-file.

ISSN: 0098-1133 (ISSN print).

DOCUMENT TYPE: Patent LANGUAGE: English

ENTRY DATE: Entered STN: 6 Feb 2003

Last Updated on STN: 6 Feb 2003

AB Compounds of the formula I, ##STR1## in which R1, R2, R3, R4, R5, and R6 have the meanings given in the description, and their physiologically acceptable salts. The compounds are suitable for use, for example, as

hypolipidemics.

L26 ANSWER 58 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

STN

ACCESSION NUMBER: 2002:618139 BIOSIS DOCUMENT NUMBER: PREV200200618139

TITLE: Effect of a newly developed specific bile acid reabsorption

inhibitor on bile acid and lipoprotein metabolism

in apoE\*3-Leiden transgenic mice.

AUTHOR(S): Groenendijk, Martine [Reprint author]; Post, Sabine M.

[Reprint author]; Schaefer, Hans-Ludwig; Kramer,

Werner; Princen, Hans M. [Reprint author]

CORPORATE SOURCE: Gaubius Laboratory, TNO-Prevention and Health, Leiden,

Netherlands

SOURCE: Hepatology, (October, 2002) Vol. 36, No. 4 Part 2, pp.

297A. print.

Meeting Info.: 53rd Annual Meeting on the Liver. BOSTON,

MA, USA. November 01-05, 2002. CODEN: HPTLD9. ISSN: 0270-9139.

DOCUMENT TYPE: Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LANGUAGE: English

ENTRY DATE: Entered STN: 4 Dec 2002

Last Updated on STN: 4 Dec 2002

L26 ANSWER 59 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

STN

ACCESSION NUMBER: 2004:7312 BIOSIS DOCUMENT NUMBER: PREV200400000600

TITLE: Prevention of cholestatic hepatitis by bile-acid-

reuptake-inhibitors in rat.

AUTHOR(S): Sauer, Peter [Reprint Author]; Kloeters-Plachky, Petra;

Kramer, Werner; Schaefer, Hans-Ludwig; Rost,

Daniel; Rudolph, Gerda; Stremmel, Wolfgang; Stiehl, Adolf

CORPORATE SOURCE: Heidelberg, Germany

SOURCE: Gastroenterology, (July 2002) Vol. 123, No. 1 Supplement,

pp. 62-63. print.

Meeting Info.: Digestive Disease Week and the 103rd Annual Meeting of the American Gastroenterological Association.

San Francisco, CA, USA. May 19-22, 2002. American

Gastroenterological Association. CODEN: GASTAB. ISSN: 0016-5085.

DOCUMENT TYPE: Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LANGUAGE: English

ENTRY DATE: Entered STN: 17 Dec 2003

Last Updated on STN: 17 Dec 2003

L26 ANSWER 60 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

STN

ACCESSION NUMBER: 2003:315418 BIOSIS DOCUMENT NUMBER: PREV200300315418

TITLE: EXPRESSION OF HUMAN Fe65 IN NEURONS AND IN VIVO.

AUTHOR(S): Santiard-Baron, D. [Reprint Author]; Delatour, B.; Clark,

A. [Reprint Author]; Schombert, B. [Reprint Author]; Touchet, N. [Reprint Author]; Bouaiche, Z. [Reprint Author]; Paul, M. F. [Reprint Author]; Duyckaerts, C.;

Pradier, L. [Reprint Author]; Benavides, J.
[Reprint Author]; Mercken, L. [Reprint Author]

CORPORATE SOURCE: Aventis Pharma, Vitry/seine, France

SOURCE: Society for Neuroscience Abstract Viewer and Itinerary

Planner, (2002) Vol. 2002, pp. Abstract No. 624.4.

http://sfn.scholarone.com. cd-rom.

Meeting Info.: 32nd Annual Meeting of the Society for Neuroscience. Orlando, Florida, USA. November 02-07, 2002.

Society for Neuroscience.

DOCUMENT TYPE: Conference; (Meeting)

Conference; (Meeting Poster)

Conference; Abstract; (Meeting Abstract)

LANGUAGE: English

ENTRY DATE: Entered STN: 9 Jul 2003

Last Updated on STN: 9 Jul 2003

AB FE65 is an adaptor protein that tightly binds to the cytoplasmic tail of the Amyloid Precursor Protein (APP). In cultured non-neuronal cells, the formation of the FE65-APP complex is a key element for the modulation of APP processing and signaling. In contrast, the function of FE65 in cultured neurons and in vivo remains to be determined. In this study, we generated double transgenic mice with neuronal expression of mutant APP and human FE65 under the control of the PDGF-B promoter (PDGF APP 695 (SDL) x PDGF FE65). Comparison of primary cultures of cortical neurons from double and single APP transgenic mice revealed that APPxFE65 neurons released more Amyloid beta-peptides in the conditioned medium than APPxWild type neurons. Our results are consistent with the data obtained in non-neuronal cells. The neuropathological status of the double

- -

transgenic mice was evaluated during their life span (up to 18 months). These mice displayed a massive glial response as abundant reactive astrocytes were observed at 9 months throughout all cortical layers (4 out of 5 animals). On the contrary, gliosis was limited and mainly evidenced into the first cortical layer in single APP transgenic mice. This neuropathological event occurred several months before the appearance of amyloid plaques, which were observed from 18 months in the single APP transgenic mice. We hypothesize that the increased release of Amyloid beta-peptides, as observed in the APPxFE65 neurons, contributes to the reactive gliosis observed in the double transgenic mice.

L26 ANSWER 61 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

STN

ACCESSION NUMBER: 2001:292955 BIOSIS DOCUMENT NUMBER: PREV200100292955

TITLE: Identification of the intestinal

cholesterol transporter.

AUTHOR(S): Kramer, W. [Reprint author]; Glombik, H. [Reprint author];

Petry, S. [Reprint author]; Heuer, H. [Reprint

author]; Corsiero, D. [Reprint author]; Girbig, F. [Reprint

author]; Weyland, C. [Reprint author]

CORPORATE SOURCE: DG Metabolic Diseases Industriepark Hoechst, Aventis Pharma

Deutschland GmbH, Gebaeude G 838, 65926, Frankfurt am Main,

Germany

SOURCE: Naunyn-Schmiedeberg's Archives of Pharmacology, (2001) Vol.

363, No. 4 Supplement, pp. R5. print.

Meeting Info.: 42nd Spring Meeting of the German Society for Experimental and Clinical Pharmacology and Toxicology. Mainz, Germany. March 13-15, 2001. German Society for Experimental and Clinical Pharmacology and Toxicology.

CODEN: NSAPCC. ISSN: 0028-1298.

DOCUMENT TYPE: Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LANGUAGE: English

ENTRY DATE: Entered STN: 20 Jun 2001

Last Updated on STN: 19 Feb 2002

L26 ANSWER 62 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

STN

ACCESSION NUMBER: 1997:512716 BIOSIS DOCUMENT NUMBER: PREV199799811919

TITLE: Topological photoaffinity labeling of the rabbit ileal

Na+/bile-salt-cotransport system.

AUTHOR(S): Kramer, Werner [Reprint author]; Wess, Guenther;

Bewersdorf, Ulrike; Corsiero, Daniel; Girbig, Frank; Weyland, Claudia; Stengelin, Siegfried; Enhsen, Alfons;

Bock, Klaus; Kleine, Horst; Le Dreau, Marie-Anne;

Schaefer, Hans-Ludwig

CORPORATE SOURCE: Hoechst Marion Roussel, DG Metabolic Dis., D-65926

Frankfurt am Main, Germany

SOURCE: European Journal of Biochemistry, (1997) Vol. 249, No. 2,

pp. 456-464.

CODEN: EJBCAI. ISSN: 0014-2956.

DOCUMENT TYPE:

Article English

LANGUAGE: ENTRY DATE:

Entered STN: 10 Dec 1997

Last Updated on STN: 10 Dec 1997

AB For the investigation of the topology of the rabbit ileal

Na+/bile-salt-cotransport system, composed of a 93-kDa integral membrane

protein and a peripheral 14-kDa bile-acid-binding protein (ILBP), we have synthesized photolabile dimeric bile-salt-transport inhibitors (photoblockers), G1-X-G2, where two bile acid moieties (G1 and G2) are tethered together via a spacer, X, and where one of the two bile acid moieties carries a photoactivatable group. These photoblockers specifically interact with the ileal Na+/ bile-salt-cotransport system as demonstrated by a concentration-dependent inhibition of (3H) cholyltaurine uptake by rabbit ileal brush-border membrane vesicles and by inhibition of photolabeling of the 93-kDa and 14-kDa bile-salt-binding proteins by 7,7-azo and 3,3-azo derivatives of cholyltaurine. Ileal bile-salt uptake was specifically inhibited by the photoblockers, which were not taken up themselves by the small intestine as demonstrated by in vivo ileal perfusion. Dependent on the photoblocker used several polypeptides in the molecular-mass range of 14-130 kDa were labeled. The cytoplasmically attached 14-kDa ILBP was significantly labeled only by inhibitors that are photoactivatable in bile acid moiety G1, suggesting that during binding and translocation of a bile-salt molecule by the ileal bile-salt-transport system the steroid nucleus gets access to the cytoplasmic site of the ileal brush-border membrane first. Photoaffinity labeling in the frozen state with the transportable 3,3-azo and 7,7-azo derivatives of cholyltaurine revealed a time-dependent increase in the extent of labeling of the 14-kDa and 93-kDa proteins, suggesting a labeling of these proteins from the cytoplasmic site of the ileal brush-border membrane. By photoaffinity labeling in the frozen state with the various photoblockers time-dependent changes in the extent of photoaffinity labeling of bile-salt-binding proteins were observed, demonstrating the possibility of topological analysis of the rabbit ileal Na+/bile-salt-cotransport system.

L26 ANSWER 63 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on STN

ACCESSION NUMBER:

1997:197782 BIOSIS

DOCUMENT NUMBER:

PREV199799496985

TITLE:

Synthesis and sar of trimeric bile acid reabsorption

inhibitors: A new approach to lower

cholesterol.

AUTHOR (S):

SOURCE:

Glombik, H.; Baringhaus, K.-H.; Boeger, G.; Enhsen, A.;

Falk, E.; Friedrich, M.; Hoffmann, A.; Kramer, W.;

Schaefer, H. L.; Stengelin, S.; Wess, G.

CORPORATE SOURCE:

HMR TA Metabolism Research D-65926 Frankfurt, Germany

Abstracts of Papers American Chemical Society, (1997) Vol.

213, No. 1-3, pp. MEDI 108.

Meeting Info.: 213th National Meeting of the American Chemical Society. San Francisco, California, USA. April

13-17, 1997.

CODEN: ACSRAL. ISSN: 0065-7727.

DOCUMENT TYPE:

Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LANGUAGE:

English

ENTRY DATE:

Entered STN: 2 May 1997

Last Updated on STN: 2 May 1997

L26 ANSWER 64 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

STN ACCESSION NUMBER:

1996:44690 BIOSIS

DOCUMENT NUMBER:

PREV199698616825

TITLE:

The ileal bile acid transporter: Molecular structure and

specific inhibitors.

AUTHOR (S):

Kramer, W. [Reprint author]; Wess, G.; Baringhaus, K.-H.;

Boeger, G.; Enhsen, A.; Falk, E.; Friedrich, M.; Glombik,

H.; Hoffmann, A.; Neckermann, G.; Pittius, C.;

Schaefer, H.-L.; Urmann, M.

CORPORATE SOURCE: Hoechst Aktiengesellschaft, D-65926 Frankfurt am Main,

Germany

2 My 2 1 1

SOURCE: Biological Chemistry Hoppe-Seyler, (1995) Vol. 376, No.

SPEC. SUPPL., pp. S70.

Meeting Info.: 120th Conference of the Gesellschaft fuer Biologische Chemie: Cell Biology and Molecular Basis of Liver Transport. Rottach-Egern, Germany. May 10-13, 1995.

CODEN: BCHSEI. ISSN: 0177-3593.

DOCUMENT TYPE: Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LANGUAGE: English

ENTRY DATE: Entered STN: 2 Feb 1996

Last Updated on STN: 3 Feb 1996

L26 ANSWER 65 OF 65 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

STN

ACCESSION NUMBER: 19

1987:272087 BIOSIS

DOCUMENT NUMBER:

PREV198784013126; BA84:13126

TITLE:

EFFECTS OF GLIADIN PEPTIDES B1-B4 IN CELIAC DISEASE I.

ORGAN CULTURE STUDIES.

AUTHOR (S):

STALLMACH A [Reprint author]; BELITZ H-D; GELLERMANN B;

SCHAEFER H; WIESER H; STERN M

CORPORATE SOURCE:

UNIV-KINDERKLINIK, MARTINISTR 52, D-2000 HAMBURG, FRG

SOURCE:

Journal of Pediatric Gastroenterology and Nutrition, (1987)

Vol. 6, No. 3, pp. 335-340. CODEN: JPGND6. ISSN: 0277-2116.

DOCUMENT TYPE:

Article

FILE SEGMENT:

ENGLISH

LANGUAGE: ENTRY DATE:

Entered STN: 19 Jun 1987

Last Updated on STN: 19 Jun 1987

Small intestinal organ culture was used as an in vitro system to study the enterotoxic effects of gliadin peptides. Measurement of enterocyte height proved to be a realiable and reproducible way of assessing mucosal change during organ culture. Enterocyte height decreases nonspecifically in normal cultured mucosal, whereas the height of enterocytes of celiac mucosa increases in vitro in controls. All the gliadin peptide fractions (B1, B2, B3, B4) that had been prepared by peptide-trypsin hydrolysis, ultrafiltration, and gel chromatography, equally inhibited the morphological increase of enterocyte height normally observed without gliadin in untreated celiac mucosa. Electrophoretic studies and amino acid analysis of B1-B4 revealed similarity between gliadin fractions with quantitative differences in molecular weight distribution of the peptide components. Our studies suggest that organ culture assessed by morphometry is a suitable model for the investigation of toxic peptides of gliadin in celiac disease. In the future, pure gliadin peptides will have to be examined.